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STL

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ANALYTICAL REPORT

PROJECT NO. 100.58.19

EMD/OHIO

Lot #: A4I270179

Angela Hurley

The Payne Firm, Inc.
11231 Cornell Park Drive
Cincinnati, OH 45242

SEVERN TRENT LABORATORIES, INC.


Roger K. Toth
Project Manager

October 5, 2004

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CASE NARRATIVE

CASE NARRATIVE

A4I270179

The following report contains the analytical results for one water sample and one quality control sample submitted to STL North Canton by The Payne Firm, Inc. from the EMD/Ohio Site, project number 100.58.19. The samples were received September 25, 2004, according to documented sample acceptance procedures.

STL utilizes USEPA approved methods in all analytical work. The samples presented in this report were analyzed for the parameter(s) listed on the analytical methods summary page in accordance with the method(s) indicated. Preliminary results were provided to Angela Hurley and Kevin Kallini on September 29, 2004, and October 03, 2004. A summary of QC data for these analyses is included at the back of the report.

STL North Canton attests to the validity of the laboratory data generated by STL facilities reported herein. All analyses performed by STL facilities were done using established laboratory SOPs that incorporate QA/QC procedures described in the applicable methods. STL's operations groups have reviewed the data for compliance with the laboratory QA/QC plan, and data have been found to be compliant with laboratory protocols unless otherwise noted below.

The test results in this report meet all NELAP requirements for parameters for which accreditation is required or available. Any exceptions to NELAP requirements are noted in this report. Pursuant to NELAP, this report may not be reproduced, except in full, without the written approval of the laboratory.

If you have any questions, please call the Project Manager, Roger K. Toth, at 330-497-9396.

This report is sequentially paginated. The final page of the report is labeled as "END OF REPORT."

SUPPLEMENTAL QC INFORMATION

SAMPLE RECEIVING

The temperature of the cooler upon sample receipt was 2.6°C.

CASE NARRATIVE (continued)

GC/MS VOLATILES

The sample(s) that contained concentrations of target analyte(s) at a reportable level in the associated Method Blank(s) were flagged with "B". All target analytes in the Method Blank must be below the reporting limit (RL) or the associated sample(s) must be ND with the exception of common laboratory contaminants.

The sample(s) that contain results between the MDL and the RL were flagged with "J". There is a possibility of false positive or mis-identification at these quantitation levels. In analytical methods requiring confirmation of the analyte reported, confirmation was performed only down to the standard reporting limit (SRL). The acceptance criteria for QC samples may not be met at these quantitation levels.

The matrix spike/matrix spike duplicate(s) for batch(es) 4273268 had recoveries outside acceptance limits. However, since the associated method blank(s) and laboratory control sample(s) were in control, no corrective action was necessary.

QUALITY CONTROL ELEMENTS OF SW-846 METHODS

STL North Canton conducts a quality assurance/quality control (QA/QC) program designed to provide scientifically valid and legally defensible data. Toward this end, several types of quality control indicators are incorporated into the QA/QC program, which is described in detail in QA Policy, QA-003. These indicators are introduced into the sample testing process to provide a mechanism for the assessment of the analytical data.

QC BATCH

Environmental samples are taken through the testing process in groups called **QUALITY CONTROL BATCHES** (QC batches). A QC batch contains up to twenty environmental samples of a similar matrix (water, soil) that are processed using the same reagents and standards. STL North Canton requires that each environmental sample be associated with a QC batch.

Several quality control samples are included in each QC batch and are processed identically to the twenty environmental samples. These QC samples include a **METHOD BLANK** (MB), a **LABORATORY CONTROL SAMPLE** (LCS) and, where appropriate, a **MATRIX SPIKE/MATRIX SPIKE DUPLICATE** (MS/MSD) pair or a **MATRIX SPIKE/SAMPLE DUPLICATE** (MS/DU) pair. If there is insufficient sample to perform an MS/MSD or an MS/DU, then a **LABORATORY CONTROL SAMPLE DUPLICATE** (LCSD) is included in the QC batch.

LABORATORY CONTROL SAMPLE

The Laboratory Control Sample is a QC sample that is created by adding known concentrations of a full or partial set of target analytes to a matrix similar to that of the environmental samples in the QC batch. The LCS analyte recovery results are used to monitor the analytical process and provide evidence that the laboratory is performing the method within acceptable guidelines. All control analytes indicated by a bold type in the LCS must meet acceptance criteria. Failure to meet the established recovery guidelines requires the repreparation and reanalysis of all samples in the QC batch. The only exception is that if the LCS recoveries are biased high and the associated sample is ND (non-detected) for the parameter(s) of interest, the batch is acceptable.

At times, a Laboratory Control Sample Duplicate (LCSD) is also included in the QC batch. An LCSD is a QC sample that is created and handled identically to the LCS. Analyte recovery data from the LCSD is assessed in the same way as that of the LCS. The LCSD recoveries, together with the LCS recoveries, are used to determine the reproducibility (precision) of the analytical system. Precision data are expressed as relative percent differences (RPDs). If the RPD fails for an LCS/LCSD and yet the recoveries are within acceptance criteria, the batch is still acceptable.

METHOD BLANK

The Method Blank is a QC sample consisting of all the reagents used in analyzing the environmental samples contained in the QC batch. Method Blank results are used to determine if interference or contamination in the analytical system could lead to the reporting of false positive data or elevated analyte concentrations. All target analytes must be below the reporting limits (RL) or the associated sample(s) must be ND except under the following circumstances:

- Common organic contaminants may be present at concentrations up to 5 times the reporting limits. Common metals contaminants may be present at concentrations up to 2 times the reporting limit, or the reported blank concentration must be twenty fold less than the concentration reported in the associated environmental samples. (See common laboratory contaminants listed below.)

<u>Volatile (GC or GC/MS)</u>	<u>Semivolatile (GC/MS)</u>	<u>Metals</u>
Methylene chloride	Phthalate Esters	Copper
Acetone		Iron
2-Butanone		Zinc
		Lead*

- *for analyses run on TJA Trace ICP, ICPMS or GFAA only*

QUALITY CONTROL ELEMENTS OF SW-846 METHODS

(Continued)

- Organic blanks will be accepted if compounds detected in the blank are present in the associated samples at levels 10 times the blank level. Inorganic blanks will be accepted if elements detected in the blank are present in the associated samples at 20 times the blank level.
- Blanks will be accepted if the compounds/elements detected are not present in any of the associated environmental samples.

Failure to meet these Method Blank criteria requires the repreparation and reanalysis of all samples in the QC batch.

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

A Matrix Spike and a Matrix Spike Duplicate are a pair of environmental samples to which known concentrations of a full or partial set of target analytes are added. The MS/MSD results are determined in the same manner as the results of the environmental sample used to prepare the MS/MSD. The analyte recoveries and the relative percent differences (RPDs) of the recoveries are calculated and used to evaluate the effect of the sample matrix on the analytical results. Due to the potential variability of the matrix of each sample, the MS/MSD results may not have an immediate bearing on any samples except the one spiked; therefore, the associated batch MS/MSD may not reflect the same compounds as the samples contained in the analytical report. When these MS/MSD results fail to meet acceptance criteria, the data is evaluated. If the LCS is within acceptance criteria, the batch is considered acceptable. The acceptance criteria do not apply to samples that are diluted for organics if the native sample amount is 4x the concentration of the spike.

For certain methods, a Matrix Spike/Sample Duplicate (MS/DU) may be included in the QC batch in place of the MS/MSD. For the parameters (i.e. pH, ignitability) where it is not possible to prepare a spiked sample, a Sample Duplicate may be included in the QC batch. However, a Sample Duplicate is less likely to provide usable precision statistics depending on the likelihood of finding concentrations below the standard reporting limit. When the Sample Duplicate result fails to meet acceptance criteria, the data is evaluated.

SURROGATE COMPOUNDS

In addition to these batch-related QC indicators, each organic environmental and QC sample is spiked with surrogate compounds. Surrogates are organic chemicals that behave similarly to the analytes of interest and that are rarely present in the environment. Surrogate recoveries are used to monitor the individual performance of a sample in the analytical system.

If surrogate recoveries are biased high in the LCS, LCSD, or the Method Blank, and the associated sample(s) are ND, the batch is acceptable. Otherwise, if the LCS, LCSD, or Method Blank surrogate(s) fail to meet recovery criteria, the entire sample batch is reprepped and reanalyzed. If the surrogate recoveries are outside criteria for environmental samples, the samples will be reprepped and reanalyzed unless there is objective evidence of matrix interference or if the sample dilution is greater than the threshold outlined in the associated method SOP.

For the GC/MS BNA methods, the surrogate criterion is that two of the three surrogates for each fraction must meet acceptance criteria. The third surrogate must have a recovery of ten percent or greater.

For the Pesticide, PCB, and PAH methods, the surrogate criterion is that one of two surrogate compounds must meet acceptance criteria.

STL North Canton Certifications and Approvals:

*California (#01144CA), Connecticut (#PH-0590), Florida (#E87225),
Illinois (#100439), Kansas (#E10336), Louisiana (#04112), Maryland (#272), Minnesota (#39-999-348), New Jersey
(#OH001), New York (#10975), Ohio (#6090), OhioVAP (#CL0024), Rhode Island (#237), South Carolina (#92007001,
#92007002, #92007003), Tennessee (#02903), Utah (#QUAN9), West Virginia (#210), Wisconsin (#999518190), NAVY,
ARMY, USDA Soil Permit, ACIL Seal of Excellence – Participating Lab Status Award (#82)*



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***EXECUTIVE
SUMMARY***

EXECUTIVE SUMMARY - Detection Highlights

AAT270179

<u>PARAMETER</u>	<u>RESULT</u>	<u>REPORTING LIMIT</u>	<u>UNITS</u>	<u>ANALYTICAL METHOD</u>
VE539/12.5-17.5/092404 09/24/04 09:58 001				
Chloromethane	0.42 J,B	1.0	ug/L	SW846 8260B
Trichloroethene	3.5	1.0	ug/L	SW846 8260B
TRIP BLANK/092404 09/24/04 002				
Acetone	5.7 J	10	ug/L	SW846 8260B
2-Butanone	2.5 J	10	ug/L	SW846 8260B
Chloromethane	0.42 J,B	1.0	ug/L	SW846 8260B
Methylene chloride	7.2	1.0	ug/L	SW846 8260B
Toluene	0.19 J	1.0	ug/L	SW846 8260B

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METHOD SUMMARY

ANALYTICAL METHODS SUMMARY

A4I270179

<u>PARAMETER</u>	<u>ANALYTICAL METHOD</u>
Volatile Organics by GC/MS	SW846 8260B

References:

SW846 "Test Methods for Evaluating Solid Waste, Physical/Chemical Methods", Third Edition, November 1986 and its updates.

SAMPLE SUMMARY

SAMPLE SUMMARY

A4I270179

WO #	SAMPLE#	CLIENT SAMPLE ID	SAMPLED DATE	SAMP TIME
GQ67F	001	VE539/12.5-17.5/092404	09/24/04	09:58
GQ67G	002	TRIP BLANK/092404	09/24/04	

NOTE(S) :

- The analytical results of the samples listed above are presented on the following pages.
- All calculations are performed before rounding to avoid round-off errors in calculated results.
- Results noted as "ND" were not detected at or above the stated limit.
- This report must not be reproduced, except in full, without the written approval of the laboratory.
- Results for the following parameters are never reported on a dry weight basis: color, corrosivity, density, flashpoint, ignitability, layers, odor, paint filter test, pH, porosity pressure, reactivity, redox potential, specific gravity, spot tests, solids, solubility, temperature, viscosity, and weight.

***SHIPPING
AND
RECEIVING DOCUMENTS***

**Chain of
Custody Record**

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STL**

STL-4124 (0901)

Client

The Payne Firm, Inc.

Address

11231 Cornell Park Dr.

City

Cinci

State

OH

Zip Code

45242

Project Name and Location (State)
Contract/Purchase Order/Quote No.

EMD/Ohio

Contract/Purchase Order/Quote No.
Carrier/Waybill Number

100-58-19

Sample I.D. No. and Description
(Containers for each sample may be combined on one line)

VE539/12.5-17.5/092404

9/24/04 0953

Trip Blank / 092404

9/24/04 —

Date

9/24/04

Time

1112

Matrix

Air

Aqueous

Sed.

Soil

Unpres.

H₂SO₄

HNO₃

HCl

NaOH

ZnAc/NaOH

#

3

2

Voc-8260

Containers & Preservatives

Analysis (Attach list if more space is needed)

Special Instructions/ Conditions of Receipt

Date

9/24/04

Page

1 or 1

Possible Hazard Identification

Non-Hazard

Flammable

Skin Irritant

Poison B

Unknown

Return To Client

Disposal By Lab

Archive For _____

Months

longer than 1 month)

Turn Around Time Required

24 Hours

48 Hours

7 Days

14 Days

21 Days

Other _____

OC Requirements (Specify)

1. Received By

Devin J. Tolok

Date

9/24/04

Time

1112

2. Received By

Devin J. Tolok

Date

9/24/04

Time

1120

3. Relinquished By

Devin J. Tolok

Date

9/24/04

Time

1112

4. Received By

Devin J. Tolok

Date

9/25/04

Time

1010

Comments

Severn Trent Laboratories, Inc.
Sample Control Record

RSR280

Client:

5670

Lot #:

A41270179

Case Number/SDG:

100.58.19

Storage Location:

MS

Laboratory Sample I.D.	Transferred By	Date	Entered	Removed	Reason	Date Returned
GQ67F	STILLERJ	9/25/04	Yes		Storage	
GQ67G	STILLERJ	9/25/04	Yes		Storage	

STL Cooler Receipt Form/Narrative

North Canton Facility

Lot Number: A4E1270F79Client: The Paville Firm

Project:

Cooler Received on: 1/25/04Opened on: 9/27/04

Quote#:

by: STL Tech
(Signature)Fedx Client Drop Off UPS DHL FAS Other: US cargo
STL Cooler No# S07 Foam Box Client Cooler Other

1. Were custody seals on the outside of the cooler? Yes No Intact? Yes No NA
If YES, Quantity 1
Were the custody seals signed and dated?
2. Shipper's packing slip attached to this form?
3. Did custody papers accompany the samples? Yes No
4. Did you sign the custody papers in the appropriate place?
5. Packing material used: Bubble Wrap Foam None
6. Cooler temperature upon receipt: -20°C (see back of form for multiple coolers/temp)
- METHOD: Temp Vial Coolant & Sample Against Bottles
COOLANT: Wet Ice Blue Ice Dry Ice Water
- Yes No NA
Yes No NA
Relinquished by client? Yes No
Yes No
Other: _____
- IR ICE/H₂O Slurry
None
Yes No
Yes No
Yes No NA
Yes No
Yes No NA
Yes No
7. Did all bottles arrive in good condition (Unbroken)?
8. Could all bottle labels and/or tags be reconciled with the COC?
9. Were samples at the correct pH? (record below/on back)
10. Were correct bottles used for the tests indicated?
11. Were air bubbles >6 mm in any VOA vials?
12. Sufficient quantity received to perform indicated analyses?

Contacted PM _____ Date: _____ by: _____ via Voice Mail Verbal Other
Concerning: _____

<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
I. CHAIN OF CUSTODY		
The following discrepancies occurred: 		

2. SAMPLE CONDITION		
Sample(s) _____	were received after the recommended holding time had expired.	
Sample(s) _____	were received in a broken container.	

3. SAMPLE PRESERVATION		
Sample(s) _____	were further preserved in sample receiving to meet recommended pH level(s). Nitric Acid Lot #052804-HNO ₃ ; Sulfuric Acid Lot # 011-504-H ₂ SO ₄ ; Sodium Hydroxide Lot # -082404-NaOH; Hydrochloric Acid Lot # 100902-HCl; Sodium Hydroxide and Zinc Acetate Lot # 071604-CH ₃ COO ₂ ZN/NaOH	
Sample(s) _____	were received with bubble > 6 mm in diameter (cc: PM)	

4. Other (see below or back)			
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>
<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>

Client ID	pH	Date	Initials

**STL Cooler Receipt Form/Narrative
North Canton Facility**

Discrepancies Cont.



GCMS VOLATILE DATA

QC SUMMARY DATA

SW846 8260B SURROGATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Lot #: A4I270179

Extraction: XXI25QK01

	CLIENT ID.	SRG01	SRG02	SRG03	SRG04	TOT OUT
01	INTRA-LAB QC	109	90	79	114	00
02	VE539/12.5-17.5/092404	113	88	81	116	00
03	TRIP BLANK/092404	113	89	83	118	00
04	METHOD BLK. GRDLR1AA	107	89	86	110	00
05	LCS GRDLR1AC	104	98	108	104	00
06	LAB MS/MSD D	109	100	112	107	00
07	LCSD GRDLR1AD	104	97	109	104	00
08	LAB MS/MSD S	110	102	114	106	00

SURROGATES

SRG01 = 1,2-Dichloroethane-d4
 SRG02 = Toluene-d8
 SRG03 = 4-Bromofluorobenzene
 SRG04 = Dibromofluoromethane

QC LIMITS

(61-128)
 (76-110)
 (74-116)
 (73-122)

- # Column to be used to flag recovery values
- * Values outside of required QC Limits
- D System monitoring Compound diluted out

FORM II

SW846 8260B CHECK SAMPLE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Lot #: A4I290000

WO #: GRDLR1AC

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	% REC	QC LIMITS REC	QUAL
Chloromethane	10	7.8	78	48- 123	
Bromomethane	10	7.3	73	64- 129	
Vinyl chloride	10	8.0	80	61- 120	
Chloroethane	10	9.2	92	66- 126	
Methylene chloride	10	13	133*	78- 118	a
Acetone	10	8.9	89	22- 200	
Carbon disulfide	10	9.8	98	73- 139	
1,1-Dichloroethene	10	10	101	63- 130	
1,1-Dichloroethane	10	9.8	98	86- 123	
1,2-Dichloroethene (total)	20	19	96	82- 116	
Chloroform	10	9.9	99	84- 128	
1,2-Dichloroethane	10	10	101	79- 136	
2-Butanone	10	7.6	76	28- 237	
1,1,1-Trichloroethane	10	8.8	88	78- 140	
Carbon tetrachloride	10	9.4	94	75- 149	
Bromodichloromethane	10	10	101	87- 130	
1,2-Dichloropropane	10	10	100	82- 115	
cis-1,3-Dichloropropene	10	9.2	92	84- 130	
Trichloroethene	10	9.6	96	75- 122	
Dibromochloromethane	10	10	101	81- 138	
1,1,2-Trichloroethane	10	9.8	98	83- 122	
Benzene	10	9.9	99	80- 116	
trans-1,3-Dichloropropene	10	8.5	85	84- 130	
Bromoform	10	11	106	76- 150	
4-Methyl-2-pentanone	10	11	107	78- 141	
2-Hexanone	10	8.1	81	35- 200	
Tetrachloroethene	10	9.6	96	88- 113	
1,1,2,2-Tetrachloroethane	10	11	113	85- 118	
Toluene	10	9.5	95	74- 119	
Chlorobenzene	10	10	102	76- 117	
Ethylbenzene	10	10	100	86- 116	

(Continued on next page)

SW846 8260B CHECK SAMPLE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Lot #: A4I290000

WO #: GRDLR1AC

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	% REC	QC LIMITS REC	QUAL
Styrene	10	11	110	85- 117	
Xylenes (total)	30	32	106	87- 116	
cis-1,2-Dichloroethene	10	9.4	94	85- 113	
trans-1,2-Dichloroethene	10	9.7	97	79- 120	
Dichlorodifluoromethane	10	6.4	64*	70- 130	a
Trichlorofluoromethane	10	8.0	80	70- 130	
1,1,2-Trichloro-1,2,2-tri	10	12	124	70- 130	
Methyl acetate	10	9.5	95	70- 130	
Methyl tert-butyl ether (10	8.6	86	70- 130	
Cyclohexane	10	8.8	88	70- 130	
Methylcyclohexane	10	8.7	87	70- 130	
1,2-Dibromoethane	10	9.8	98	70- 130	
Isopropylbenzene	10	11	110	70- 130	
1,3-Dichlorobenzene	10	9.8	98	70- 130	
1,4-Dichlorobenzene	10	10	103	70- 130	
1,2-Dichlorobenzene	10	9.8	98	70- 130	
1,2-Dibromo-3-chloropropane	10	7.6	76	70- 130	
1,2,4-Trichlorobenzene	10	3.6	36*	70- 130	a

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

* Values outside of QC limits

Spike Recovery: 3 out of 49 outside limits

COMMENTS:

SW846 8260B CHECK SAMPLE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Lot #: A4I290000

WO #: GRDLR1AD

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	% REC	QC LIMITS REC	QUAL
Chloromethane	10	8.2	82	48 - 123	
Bromomethane	10	7.8	78	64 - 129	
Vinyl chloride	10	8.2	82	61 - 120	
Chloroethane	10	9.6	96	66 - 126	
Methylene chloride	10	14	138*	78 - 118	a
Acetone	10	9.1	91	22 - 200	
Carbon disulfide	10	10	104	73 - 139	
1,1-Dichloroethene	10	10	103	63 - 130	
1,1-Dichloroethane	10	10	100	86 - 123	
1,2-Dichloroethene (total)	20	20	99	82 - 116	
Chloroform	10	9.7	97	84 - 128	
1,2-Dichloroethane	10	10	101	79 - 136	
2-Butanone	10	8.7	87	28 - 237	
1,1,1-Trichloroethane	10	8.7	87	78 - 140	
Carbon tetrachloride	10	9.6	96	75 - 149	
Bromodichloromethane	10	9.9	99	87 - 130	
1,2-Dichloropropane	10	9.6	96	82 - 115	
cis-1,3-Dichloropropene	10	9.2	92	84 - 130	
Trichloroethene	10	9.6	96	75 - 122	
Dibromochloromethane	10	9.9	99	81 - 138	
1,1,2-Trichloroethane	10	9.9	99	83 - 122	
Benzene	10	9.7	97	80 - 116	
trans-1,3-Dichloropropene	10	8.4	84	84 - 130	
Bromoform	10	11	110	76 - 150	
4-Methyl-2-pentanone	10	12	117	78 - 141	
2-Hexanone	10	8.6	86	35 - 200	
Tetrachloroethene	10	9.8	98	88 - 113	
1,1,2,2-Tetrachloroethane	10	12	116	85 - 118	
Toluene	10	9.4	94	74 - 119	
Chlorobenzene	10	10	100	76 - 117	
Ethylbenzene	10	10	100	86 - 116	

(Continued on next page)

SW846 8260B CHECK SAMPLE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Lot #: A4I290000

WO #: GRDLR1AD

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	% REC	QC LIMITS REC	QUAL
Styrene	10	11	108	85- 117	
Xylenes (total)	30	31	105	87- 116	
cis-1,2-Dichloroethene	10	9.5	95	85- 113	
trans-1,2-Dichloroethene	10	10	102	79- 120	
Dichlorodifluoromethane	10	6.9	69*	70- 130	a
Trichlorofluoromethane	10	8.2	82	70- 130	
1,1,2-Trichloro-1,2,2-tri	10	13	126	70- 130	
Methyl acetate	10	10	100	70- 130	
Methyl tert-butyl ether (10	8.3	83	70- 130	
Cyclohexane	10	9.2	92	70- 130	
Methylcyclohexane	10	9.0	90	70- 130	
1,2-Dibromoethane	10	10	100	70- 130	
Isopropylbenzene	10	11	111	70- 130	
1,3-Dichlorobenzene	10	9.4	94	70- 130	
1,4-Dichlorobenzene	10	10	100	70- 130	
1,2-Dichlorobenzene	10	9.4	94	70- 130	
1,2-Dibromo-3-chloropropane	10	7.8	78	70- 130	
1,2,4-Trichlorobenzene	10	4.0	40*	70- 130	a

NOTES(S) :

a Spiked analyte recovery is outside stated control limits.

* Values outside of QC limits

Spike Recovery: 3 out of 49 outside limits

COMMENTS:

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Matrix Spike ID: LAB MS/MSD

Lot #: A4I180226

WO #: GQKRM1AC

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	MS CONCENT. (ug/L)	MS % REC	LIMITS REC	QUAL
1,1-Dichloroethene	710	ND	780	109	62- 130	
Chloromethane	710	24	590	79	40- 137	
Bromomethane	710	ND	510	72	55- 145	
Vinyl chloride	710	ND	600	83*	88- 126	a
Chloroethane	710	ND	730	102	59- 142	
Methylene chloride	710	160	950	111	82- 115	
Acetone	710	110	650	76	45- 128	
Carbon disulfide	710	ND	800	112	69- 138	
1,1-Dichloroethane	710	ND	740	103	88- 127	
1,2-Dichloroethene (total)	1400	910	2400	101	86- 115	
Chloroform	710	ND	730	102	83- 141	
1,2-Dichloroethane	710	ND	740	104	71- 160	
2-Butanone	710	ND	580	81	71- 123	
1,1,1-Trichloroethane	710	ND	650	91	71- 162	
Carbon tetrachloride	710	ND	760	106	63- 176	
Bromodichloromethane	710	ND	730	102	80- 146	
1,2-Dichloropropane	710	ND	710	100	87- 114	
cis-1,3-Dichloropropene	710	ND	630	89	82- 130	
Trichloroethene	710	2100	2600	75	62- 130	
Dibromochloromethane	710	ND	740	104	71- 158	
1,1,2-Trichloroethane	710	ND	720	101	86- 129	
Benzene	710	ND	740	101	78- 118	
trans-1,3-Dichloropropene	710	ND	610	85	73- 147	
Bromoform	710	ND	790	111	58- 176	
4-Methyl-2-pentanone	710	ND	780	109	82- 135	
2-Hexanone	710	ND	600	84	81- 128	
Tetrachloroethene	710	100	860	107	85- 121	
1,1,2,2-Tetrachloroethane	710	ND	830	117*	88- 116	a

(Continued on next page)

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Matrix Spike ID: LAB MS/MSD

Lot #: A4I180226

WO #: GQKRM1AC

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE CONCENT. (ug/L)	MS CONCENT. (ug/L)	MS % REC	LIMITS REC	QUAL
Toluene	710	ND	710	99	70 - 119	
Chlorobenzene	710	ND	730	102	76 - 117	
Ethylbenzene	710	ND	740	103	86 - 132	
Styrene	710	ND	800	113	83 - 120	
Xylenes (total)	2100	ND	2400	110	89 - 121	
cis-1,2-Dichloroethene	710	900	1600	97	87 - 114	
trans-1,2-Dichloroethene	710	12	760	104	85 - 116	
Dichlorodifluoromethane	710	ND	530	74	70 - 130	
Trichlorofluoromethane	710	ND	670	94	70 - 130	
1,1,2-Trichloro-1,2,2-tri	710	ND	1100	149*	70 - 130	a
Methyl acetate	710	ND	700	97	70 - 130	
Methyl tert-butyl ether (710	ND	610	86	70 - 130	
Cyclohexane	710	ND	740	103	70 - 130	
Methylcyclohexane	710	ND	740	103	70 - 130	
1,2-Dibromoethane	710	ND	730	102	70 - 130	
Isopropylbenzene	710	ND	840	117	70 - 130	
1,3-Dichlorobenzene	710	ND	670	94	70 - 130	
1,4-Dichlorobenzene	710	ND	740	104	70 - 130	
1,2-Dichlorobenzene	710	ND	670	94	70 - 130	
1,2-Dibromo-3-chloropropane	710	ND	530	74	70 - 130	
1,2,4-Trichlorobenzene	710	ND	280	39*	70 - 130	a

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

RPD: 0 out of 0 outside limits

Spike Recovery: 4 out of 49 outside limits

COMMENTS:

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc. Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Matrix Spike ID: LAB MS/MSD

Lot #: A4I180226

WO #: GQKRM1AD

BATCH: 4273268

COMPOUND	SPIKE ADDED (ug/L)	MSD CONCENT. (ug/L)	MSD % REC	MSD % RPD	QC LIMITS RPD	QC LIMITS REC	QUAL
2-Hexanone	710	550	77*	7.5	30	81- 128	a
Tetrachloroethene	710	800	98	7.5	30	85- 121	
1,1,2,2-Tetrachloroethane	710	810	114	2.7	30	88- 116	
Toluene	710	680	95	3.4	20	70- 119	
Chlorobenzene	710	720	101	1.6	20	76- 117	
Ethylbenzene	710	710	99	3.9	30	86- 132	
Styrene	710	770	108	4.3	30	83- 120	
Xylenes (total)	2100	2200	105	4.6	30	89- 121	
cis-1,2-Dichloroethene	710	1500	88	4.0	30	87- 114	
trans-1,2-Dichloroethene	710	720	100	4.3	30	85- 116	
Dichlorodifluoromethane	710	500	71	4.9	30	70- 130	
Trichlorofluoromethane	710	600	84	11	30	70- 130	
1,1,2-Trichloro-1,2,2-tri	710	890	125	18	30	70- 130	
Methyl acetate	710	670	94	4.0	30	70- 130	
Methyl tert-butyl ether (710	610	85	0.97	30	70- 130	
Cyclohexane	710	660	92	11	30	70- 130	
Methylcyclohexane	710	660	92	11	30	70- 130	
1,2-Dibromoethane	710	700	98	3.8	30	70- 130	
Isopropylbenzene	710	780	109	6.7	30	70- 130	
1,3-Dichlorobenzene	710	670	94	0.39	30	70- 130	
1,4-Dichlorobenzene	710	730	102	1.9	30	70- 130	
1,2-Dichlorobenzene	710	680	95	1.1	30	70- 130	
1,2-Dibromo-3-chloropropa	710	510	72	3.0	30	70- 130	
1,2,4-Trichlorobenzene	710	260	37*	7.5	30	70- 130	a
1,1-Dichloroethene	710	740	104	4.6	20	62- 130	
Chloromethane	710	550	74	6.8	39	40- 137	
Bromomethane	710	550	77	7.4	30	55- 145	
Vinyl chloride	710	610	84*	1.2	30	88- 126	a

(Continued on next page)

SW846 8260B MATRIX SPIKE/MATRIX SPIKE DUPLICATE RECOVERY

Lab Name: Severn Trent Laboratories, Inc.

Client: PAYNE FIRM INC.

Lab Code: STLCAN

SDG No: 4I27179

Matrix Spike ID: LAB MS/MSD

Lot #: A4II180226

WO #: GQKRM1AD

BATCH: 4273268

COMPOUND	SPIKE ADDED	MSD CONCENT.	MSD %	MSD %	QC LIMITS		QUAL
			REC	RPD	RPD	REC	
Chloroethane	710	710	99	3.2	-	30	59- 142
Methylene chloride	710	930	108	2.2	-	30	82- 115
Acetone	710	610	70	6.2	-	30	45- 128
Carbon disulfide	710	730	102	9.2	-	41	69- 138
1,1-Dichloroethane	710	720	101	2.4	-	30	88- 127
1,2-Dichloroethene (total)	1400	2300	94	4.1	-	30	86- 115
Chloroform	710	700	98	4.2	-	30	83- 141
1,2-Dichloroethane	710	710	99	4.7	-	30	71- 160
2-Butanone	710	580	82	1.3	-	30	71- 123
1,1,1-Trichloroethane	710	620	86	5.0	-	30	71- 162
Carbon tetrachloride	710	700	98	7.4	-	30	63- 176
Bromodichloromethane	710	710	99	2.4	-	30	80- 146
1,2-Dichloropropane	710	710	100	0.0	-	30	87- 114
cis-1,3-Dichloropropene	710	620	87	2.3	-	30	82- 130
Trichloroethene	710	2500	53*	6.2	-	20	62- 130 a
Dibromochloromethane	710	700	98	6.1	-	30	71- 158
1,1,2-Trichloroethane	710	690	97	4.8	-	30	86- 129
Benzene	710	710	97	3.8	-	20	78- 118
trans-1,3-Dichloropropene	710	570	79	7.0	-	30	73- 147
Bromoform	710	720	101	9.7	-	30	58- 176
4-Methyl-2-pentanone	710	750	105	4.3	-	30	82- 135

NOTES (S) :

a Spiked analyte recovery is outside stated control limits.

Column to be used to flag recovery and RPD values with an asterisk
* Values outside of QC limitsRPD: 0 out of 49 outside limits
Spike Recovery: 4 out of 49 outside limits

COMMENTS:

BLANK WORKORDER NO.

SW846 8260B METHOD BLANK SUMMARY

GRDLR1AA

Lab Name: Severn Trent Laboratories, Inc.

Lab Code: STLCAN

SDG Number: 4I27179

Lab File ID: UXJ24193.

Lot Number: A4I270179

Date Analyzed: 09/29/04

Time Analyzed: 09:22

Matrix: WATER

Date Extracted: 09/29/04

GC Column: DB 624 ID: .18

Extraction Method: 5030B/8260B

Instrument ID: UX11

Level: (low/med) LOW

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, LCS, LCSD, MS , MSD:

CLIENT ID.	SAMPLE	LAB	DATE	TIME
	WORK ORDER #	FILE ID	ANALYZED	ANALYZED
01 INTRA-LAB QC	GQKRM1AA	UXJ24202.	09/29/04	12:49
02 LAB MS/MSD	GQKRM1AC S	UXJ24211.	09/29/04	16:14
03 LAB MS/MSD	GQKRM1AD D	UXJ24212.	09/29/04	16:37
04 VE539/12.5-17.5/092404	GQ67F1AA	UXJ24200.	09/29/04	12:03
05 TRIP BLANK/092404	GQ67G1AA	UXJ24201.	09/29/04	12:26
06 CHECK SAMPLE	GRDLR1AC C	UXJ24191.	09/29/04	08:37
07 DUPLICATE CHECK	GRDLR1AD L	UXJ24192.	09/29/04	09:00
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COMMENTS:

5A
VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.:

SDG No.: 4I27179

Lab File ID: BFB207

BFB Injection Date: 08/16/04

Instrument ID: A3UX11

BFB Injection Time: 1309

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	16.8
75	30.0 - 60.0% of mass 95	44.5
95	Base Peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.9
173	Less than 2.0% of mass 174	0.7 (0.8)1
174	50.0 - 100.0% of mass 95	81.3
175	5.0 - 9.0% of mass 174	6.1 (7.5)1
176	Greater than 95.0%, but less than 101.0% of mass 174	78.9 (97.1)1
177	5.0 - 9.0% of mass 176	5.0 (6.4)2

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01 VSTD040	200NG-A9IC	UXJ23209	08/16/04	1618
02 VSTD020	100NG-A9IC	UXJ23210	08/16/04	1640
03 VSTD010	50NG-A9IC	UXJ23211	08/16/04	1703
04 VSTD005	25NG-A9IC	UXJ23212	08/16/04	1726
05 VSTD002	10NG-A9IC	UXJ23213	08/16/04	1748
06 VSTD001	5NG-A9IC	UXJ23214	08/16/04	1811
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5A
VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.:

SDG No.: 4I27179

Lab File ID: BFB232

BFB Injection Date: 09/14/04

Instrument ID: A3UX11

BFB Injection Time: 1321

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	19.8
75	30.0 - 60.0% of mass 95	51.6
95	Base Peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.6
173	Less than 2.0% of mass 174	0.2 (0.3)1
174	50.0 - 100.0% of mass 95	74.6
175	5.0 - 9.0% of mass 174	5.6 (7.5)1
176	Greater than 95.0%, but less than 101.0% of mass 174	74.5 (99.9)1
177	5.0 - 9.0% of mass 176	5.2 (6.9)2

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01 VSTD040	200NG-IC	UXJ23870	09/14/04	1348
02 VSTD020	100NG-IC	UXJ23871	09/14/04	1410
03 VSTD010	50NG-IC	UXJ23872	09/14/04	1433
04 VSTD005	25NG-IC	UXJ23873	09/14/04	1457
05 VSTD002	10NG-IC	UXJ23874	09/14/04	1519
06 VSTD001	5NG-IC	UXJ23875	09/14/04	1541
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5A
VOLATILE ORGANIC GC/MS TUNING AND MASS
CALIBRATION - BROMOFLUOROBENZENE (BFB)

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.:

SDG No.: 4I27179

Lab File ID: BFB243

BFB Injection Date: 09/29/04

Instrument ID: A3UX11

BFB Injection Time: 0725

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	18.8
75	30.0 - 60.0% of mass 95	50.2
95	Base Peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.8
173	Less than 2.0% of mass 174	0.4 (0.5)1
174	50.0 - 100.0% of mass 95	75.5
175	5.0 - 9.0% of mass 174	4.9 (6.4)1
176	Greater than 95.0%, but less than 101.0% of mass 174	73.3 (97.0)1
177	5.0 - 9.0% of mass 176	4.6 (6.3)2

1-Value is % of mass 174

2-Value is % of mass 176

THIS TUNE APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01 VSTD010	50NG-CC	UXJ24189	09/29/04	0750
02 VSTD010	50NG-A9CC	UXJ24190	09/29/04	0814
03 GRDLR-CHK	GRDLR1AC	UXJ24191	09/29/04	0837
04 GRDLR-CKDUP	GRDLR1AD	UXJ24192	09/29/04	0900
05 GRDLR-BLK	GRDLR1AA	UXJ24193	09/29/04	0922
06 VE539/12.5-1	GQ67F1AA	UXJ24200	09/29/04	1203
07 TRIP BLANK/0	GQ67G1AA	UXJ24201	09/29/04	1226
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8A
VOLATILE INTERNAL STANDARD AREA SUMMARY

Lab Name: STL - NORTH CANTON

Contract:

Lab Code: STLCAN Case No.:

SAS No.:

SDG No.: 4I27179

Lab File ID (Standard): UXJ24189

Date Analyzed: 09/29/04

Instrument ID: A3UX11

Time Analyzed: 0750

Matrix: (soil/water) WATER Level: (low/med) LOW Column: (pack/cap) CAP

	IS1(CBZ) AREA #	RT	IS2 AREA #	RT	IS3(DCB) AREA #	RT
12 HOUR STD	1787660	7.68	2187253	5.04	1037377	9.90
UPPER LIMIT	3575320	8.18	4374506	5.54	2074754	10.40
LOWER LIMIT	893830	7.18	1093627	4.54	518689	9.40
EPA SAMPLE NO.						
01 GRDLR-CHK	1771165	7.68	2102753	5.04	886729	9.90
02 GRDLR-CKDUP	1767481	7.68	2090526	5.04	932881	9.90
03 GRDLR-BLK	1674688	7.68	1924922	5.04	707660	9.90
04 VE539/12.5-1	1628364	7.68	1840075	5.04	670401	9.90
05 TRIP BLANK/0	1610955	7.68	1816483	5.04	682806	9.90
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22						

IS1 (CBZ) = Chlorobenzene-d5

IS2 = Fluorobenzene

IS3 (DCB) = 1,4-Dichlorobenzene-d4

UPPER LIMIT = +100%

of internal standard area.

LOWER LIMIT = - 50%

of internal standard area.

Column used to flag internal standard area values with an asterisk.

SAMPLE DATA

PAYNE FIRM INC.

Client Sample ID: VE539/12.5-17.5/092404

GC/MS Volatiles

Lot-Sample #....: A4I270179-001 Work Order #....: GQ67F1AA Matrix.....: WG
 Date Sampled....: 09/24/04 09:58 Date Received..: 09/25/04
 Prep Date.....: 09/29/04 Analysis Date...: 09/29/04
 Prep Batch #....: 4273268
 Dilution Factor: 1 Initial Wgt/Vol: 5 mL Final Wgt/Vol.: 5 mL
 Method.....: SW846 8260B

PARAMETER	RESULT	REPORTING	
		LIMIT	UNITS
Acetone	ND	10	ug/L
Acetonitrile	ND	20	ug/L
Acrolein	ND	20	ug/L
Acrylonitrile	ND	20	ug/L
Benzene	ND	1.0	ug/L
Bromodichloromethane	ND	1.0	ug/L
Bromoform	ND	1.0	ug/L
Bromomethane	ND	1.0	ug/L
2-Butanone	ND	10	ug/L
Carbon disulfide	ND	1.0	ug/L
Carbon tetrachloride	ND	1.0	ug/L
Chlorobenzene	ND	1.0	ug/L
Chloroprene	ND	2.0	ug/L
Dibromochloromethane	ND	1.0	ug/L
Chloroethane	ND	1.0	ug/L
Chloroform	ND	1.0	ug/L
Chloromethane	0.42 J,B	1.0	ug/L
3-Chloropropene	ND	2.0	ug/L
1,2-Dibromo-3-chloropropane (DBCP)	ND	2.0	ug/L
1,2-Dibromoethane	ND	1.0	ug/L
Dibromomethane	ND	1.0	ug/L
trans-1,4-Dichloro-2-butene	ND	1.0	ug/L
1,1-Dichloroethane	ND	1.0	ug/L
1,2-Dichloroethane	ND	1.0	ug/L
cis-1,2-Dichloroethene	ND	1.0	ug/L
trans-1,2-Dichloroethene	ND	1.0	ug/L
1,1-Dichloroethene	ND	1.0	ug/L
1,2-Dichloroethene (total)	ND	2.0	ug/L
Dichlorofluoromethane	ND	2.0	ug/L
1,2-Dichloropropane	ND	1.0	ug/L
cis-1,3-Dichloropropene	ND	1.0	ug/L
trans-1,3-Dichloropropene	ND	1.0	ug/L
1,4-Dioxane	ND	50	ug/L
Ethylbenzene	ND	1.0	ug/L
Ethyl methacrylate	ND	1.0	ug/L

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: VE539/12.5-17.5/092404

GC/MS Volatiles

Lot-Sample #....: A4I270179-001 Work Order #....: GQ67F1AA Matrix.....: WG

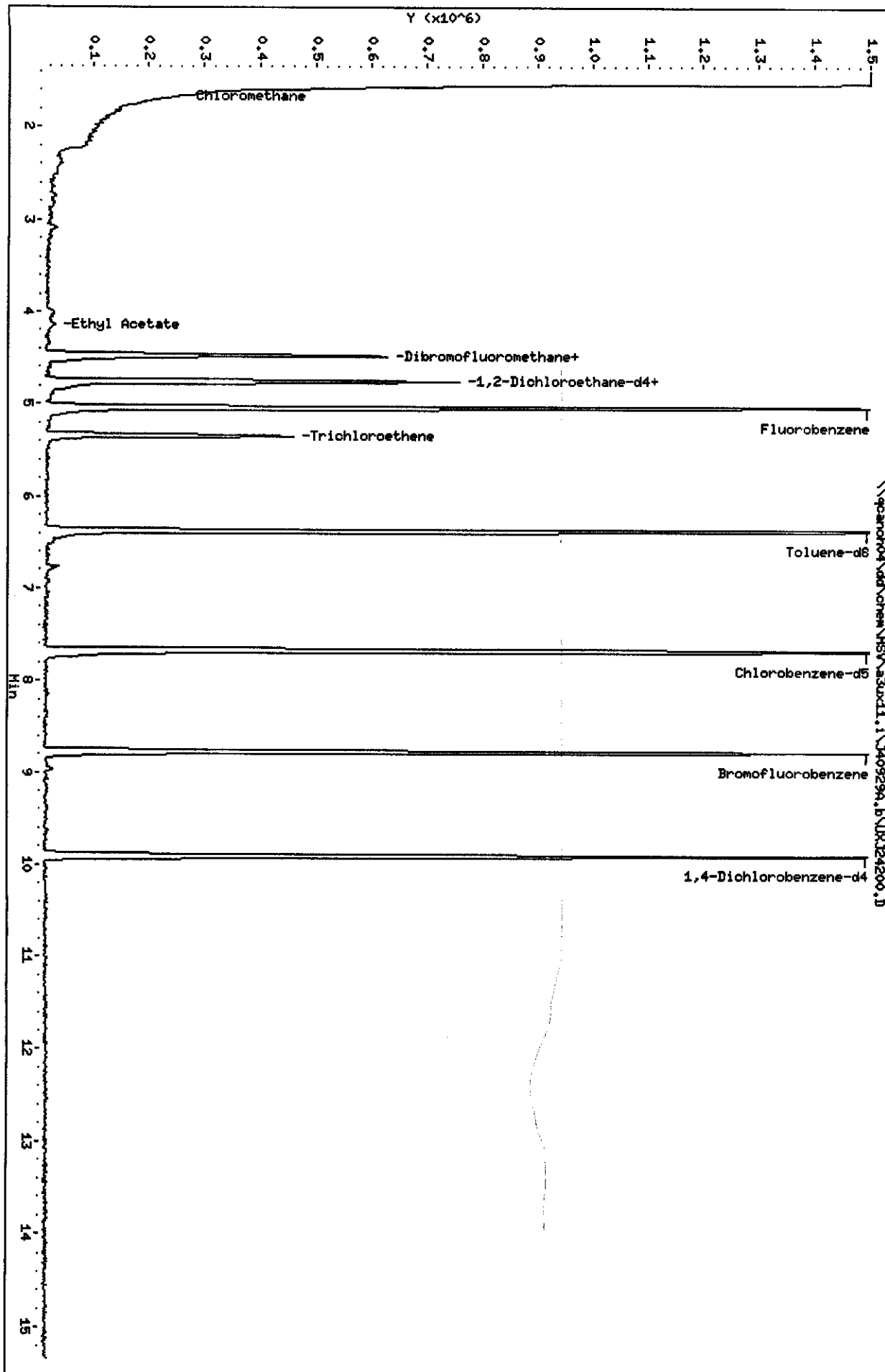
<u>PARAMETER</u>	<u>RESULT</u>	<u>REPORTING LIMIT</u>	<u>UNITS</u>
2-Hexanone	ND	10	ug/L
Iodomethane	ND	1.0	ug/L
Isobutanol	ND	50	ug/L
Methacrylonitrile	ND	2.0	ug/L
Methylene chloride	ND	1.0	ug/L
Methyl methacrylate	ND	2.0	ug/L
4-Methyl-2-pentanone	ND	10	ug/L
Propionitrile	ND	4.0	ug/L
Styrene	ND	1.0	ug/L
1,1,1,2-Tetrachloroethane	ND	1.0	ug/L
1,1,2,2-Tetrachloroethane	ND	1.0	ug/L
Tetrachloroethene	ND	1.0	ug/L
Toluene	ND	1.0	ug/L
1,1,1-Trichloroethane	ND	1.0	ug/L
1,1,2-Trichloroethane	ND	1.0	ug/L
Trichloroethene	3.5	1.0	ug/L
Trichlorofluoromethane	ND	1.0	ug/L
1,2,3-Trichloropropane	ND	1.0	ug/L
Vinyl acetate	ND	2.0	ug/L
Vinyl chloride	ND	1.0	ug/L
Xylenes (total)	ND	2.0	ug/L

<u>SURROGATE</u>	<u>PERCENT RECOVERY</u>	<u>RECOVERY LIMITS</u>
Dibromofluoromethane	116	(73 - 122)
1,2-Dichloroethane-d4	113	(61 - 128)
Toluene-d8	88	(76 - 110)
4-Bromofluorobenzene	81	(74 - 116)

NOTE(S) :

J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.



Data File: \\pcand04\dat\chen\HSV\230x1.i\\40929A.b\UX24200.D
 Date : 29-SEP-2004 12:03
 Client ID: VES3942.5-17.5/092
 Sample Info: G6751AA,5ML/5ML
 Purge Volume: 5.0
 Column phase: DB624

 Instrument: 230x1.i
 Operator: 43582
 Column diameter: 0.18
 \\pcand04\dat\chen\HSV\230x1.i\\40929A.b\UX24200.D

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24200.D
Lab Smp Id: GQ67F1AA Client Smp ID: VE539/12.5-17.5/092
Inj Date : 29-SEP-2004 12:03
Operator : 43582 Inst ID: a3ux11.i
Smp Info : GQ67F1AA, 5ML/5ML
Misc Info : J40929A, 8260LLUX11, , 43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 12
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ug/L)
*	1 Fluorobenzene	96	5.041	5.041 (1.000)	1840075	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1628364	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	670401	50.0000		
\$	4 Dibromofluoromethane	113	4.485	4.473 (0.890)	497468	58.0667	11.613	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	666205	56.5513	11.310	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	1722801	44.0391	8.808	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	671180	40.3799	8.076	
8	Dichlorodifluoromethane	85		Compound Not Detected.				
9	Chloromethane	50	1.669	1.692 (0.931)	36465	2.12303	0.4246	
10	Vinyl Chloride	62		Compound Not Detected.				
11	Bromomethane	94		Compound Not Detected.				
12	Chloroethane	64		Compound Not Detected.				
13	Trichlorofluoromethane	101		Compound Not Detected.				
15	Acrolein	56		Compound Not Detected.				
16	Acetone	43		Compound Not Detected.				
17	1,1-Dichloroethene	96		Compound Not Detected.				
18	Freon-113	151		Compound Not Detected.				

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
19 Iodomethane	142					Compound Not Detected.	
20 Carbon Disulfide	76					Compound Not Detected.	
21 Methylene Chloride	84					Compound Not Detected.	
22 Acetonitrile	41					Compound Not Detected.	
23 Acrylonitrile	53					Compound Not Detected.	
24 Methyl tert-butyl ether	73					Compound Not Detected.	
25 trans-1,2-Dichloroethene	96					Compound Not Detected.	
26 Hexane	86					Compound Not Detected.	
27 Vinyl acetate	43					Compound Not Detected.	
28 1,1-Dichloroethane	63					Compound Not Detected.	
29 tert-Butyl Alcohol	59					Compound Not Detected.	
30 2-Butanone	43					Compound Not Detected.	
M 31 1,2-Dichloroethene (total)	96					Compound Not Detected.	
32 cis-1,2-dichloroethene	96					Compound Not Detected.	
33 2,2-Dichloropropane	77					Compound Not Detected.	
34 Bromochloromethane	128					Compound Not Detected.	
35 Chloroform	83					Compound Not Detected.	
36 Tetrahydrofuran	42					Compound Not Detected.	
37 1,1,1-Trichloroethane	97	4.508	4.520 (0.894)		13024	0.96333	0.1927
38 1,1-Dichloropropene	75					Compound Not Detected.	
39 Carbon Tetrachloride	117					Compound Not Detected.	
40 1,2-Dichloroethane	62					Compound Not Detected.	
41 Benzene	78	4.816	4.816 (0.955)		35544	0.83901	0.1678
42 Trichloroethene	130	5.349	5.349 (1.061)		171287	17.7158	3.543
43 1,2-Dichloropropane	63					Compound Not Detected.	
44 1,4-Dioxane	88					Compound Not Detected.	
45 Dibromomethane	93					Compound Not Detected.	
46 Bromodichloromethane	83					Compound Not Detected.	
47 2-Chloroethyl vinyl ether	63					Compound Not Detected.	
48 cis-1,3-Dichloropropene	75					Compound Not Detected.	
49 4-Methyl-2-pentanone	43					Compound Not Detected.	
50 Toluene	91					Compound Not Detected.	
51 trans-1,3-Dichloropropene	75					Compound Not Detected.	
52 Ethyl Methacrylate	69					Compound Not Detected.	
53 1,1,2-Trichloroethane	97					Compound Not Detected.	
54 1,3-Dichloropropane	76					Compound Not Detected.	
55 Tetrachloroethene	164					Compound Not Detected.	
56 2-Hexanone	43					Compound Not Detected.	
57 Dibromochloromethane	129					Compound Not Detected.	
58 1,2-Dibromoethane	107					Compound Not Detected.	
59 Chlorobenzene	112					Compound Not Detected.	
60 1,1,1,2-Tetrachloroethane	131					Compound Not Detected.	
61 Ethylbenzene	106					Compound Not Detected.	
62 m + p-Xylene	106					Compound Not Detected.	
M 63 Xylenes (total)	106					Compound Not Detected.	
64 Xylene-o	106					Compound Not Detected.	
65 Styrene	104					Compound Not Detected.	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24200.D
 Report Date: 30-Sep-2004 08:43

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng)	FINAL (ug/L)
66 Bromoform	173					Compound Not Detected.		
67 Isopropylbenzene	105					Compound Not Detected.		
68 1,1,2,2-Tetrachloroethane	83					Compound Not Detected.		
69 1,4-Dichloro-2-butene	53					Compound Not Detected.		
70 1,2,3-Trichloropropane	110					Compound Not Detected.		
71 Bromobenzene	156					Compound Not Detected.		
72 n-Propylbenzene	120					Compound Not Detected.		
73 2-Chlorotoluene	126					Compound Not Detected.		
74 1,3,5-Trimethylbenzene	105					Compound Not Detected.		
75 4-Chlorotoluene	126					Compound Not Detected.		
76 tert-Butylbenzene	119					Compound Not Detected.		
77 1,2,4-Trimethylbenzene	105					Compound Not Detected.		
78 sec-Butylbenzene	105					Compound Not Detected.		
79 4-Isopropyltoluene	119					Compound Not Detected.		
80 1,3-Dichlorobenzene	146					Compound Not Detected.		
81 1,4-Dichlorobenzene	146					Compound Not Detected.		
82 n-Butylbenzene	91					Compound Not Detected.		
83 1,2-Dichlorobenzene	146					Compound Not Detected.		
84 1,2-Dibromo-3-chloropropane	157					Compound Not Detected.		
85 1,2,4-Trichlorobenzene	180					Compound Not Detected.		
86 Hexachlorobutadiene	225					Compound Not Detected.		
87 Naphthalene	128					Compound Not Detected.		
88 1,2,3-Trichlorobenzene	180					Compound Not Detected.		
14 Dichlorofluoromethane	67					Compound Not Detected.		
89 Ethyl Ether	59					Compound Not Detected.		
91 3-Chloropropene	76					Compound Not Detected.		
92 Isopropyl Ether	87					Compound Not Detected.		
93 2-Chloro-1,3-butadiene	53					Compound Not Detected.		
94 Propionitrile	54					Compound Not Detected.		
95 Ethyl Acetate	43	4.153	4.154	(0.824)	26303	2.91627	0.5832	
96 Methacrylonitrile	41					Compound Not Detected.		
97 Isobutanol	41					Compound Not Detected.		
99 n-Butanol	56					Compound Not Detected.		
100 Methyl Methacrylate	41					Compound Not Detected.		
101 2-Nitropropane	41					Compound Not Detected.		
103 Cyclohexanone	55					Compound Not Detected.		
98 Cyclohexane	56					Compound Not Detected.		
143 Methyl Acetate	43					Compound Not Detected.		
144 Methylcyclohexane	83					Compound Not Detected.		
141 1,3,5-Trichlorobenzene	180					Compound Not Detected.		

Data File: \\qcanch04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UXJ24200.D

Date : 29-SEP-2004 12:03

Client ID: VE539/12.5-17.5/092

Instrument: a3ux11.i

Sample Info: GQ67F1AA,5ML/5ML

Purge Volume: 5.0

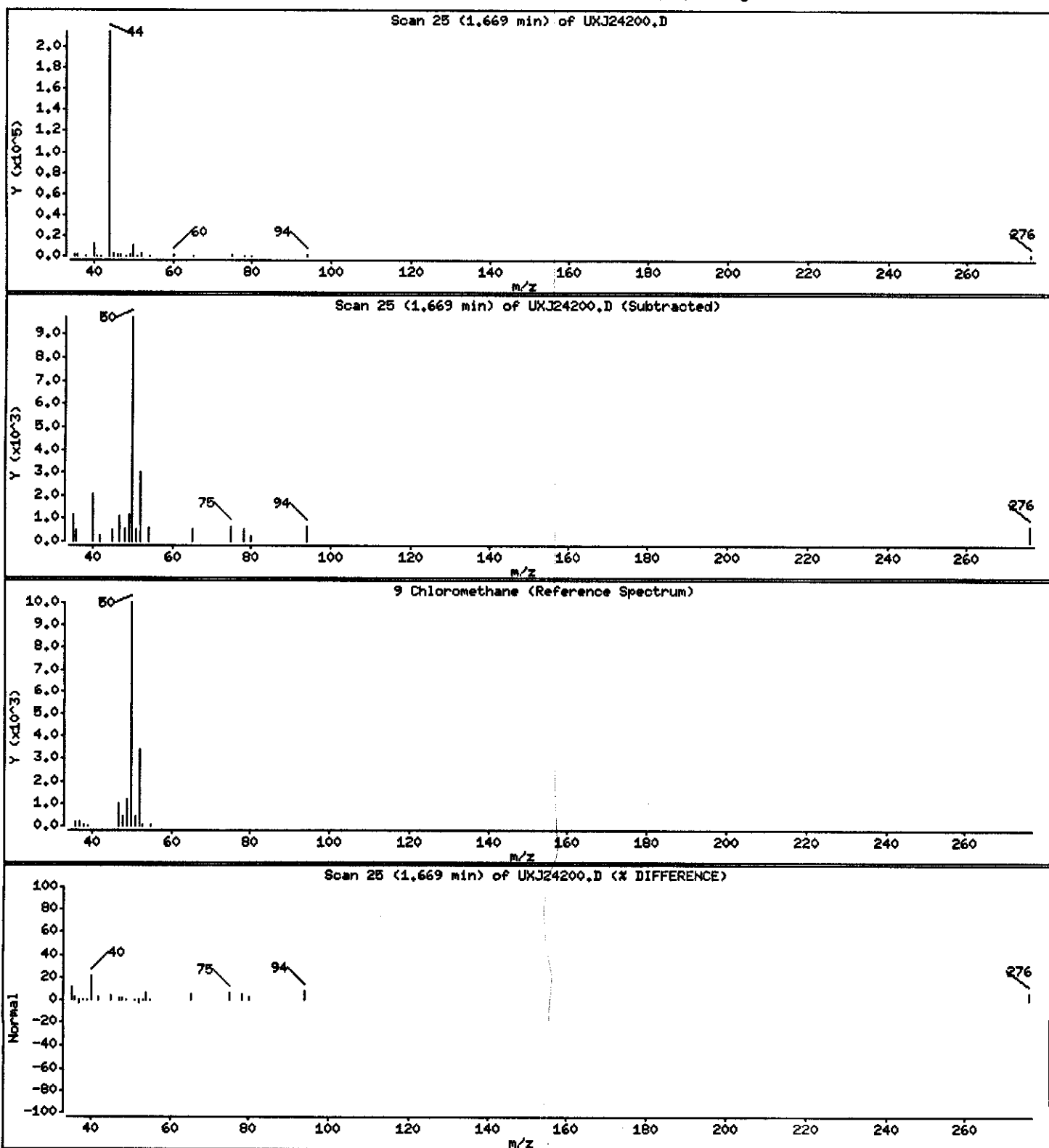
Operator: 43582

Column phase: DB624

Column diameter: 0.18

9 Chloromethane

Concentration: 0.4246 ug/L



Data File: \\qcanoh04\dd\chem\MSV\3ux11.i\J40929A.b\UXJ24200.D

Date : 29-SEP-2004 12:03

Client ID: VE539/12.5-17.5/092

Instrument: 3ux11.i

Sample Info: GQ67F1AA,5ML/5ML

Purge Volume: 5.0

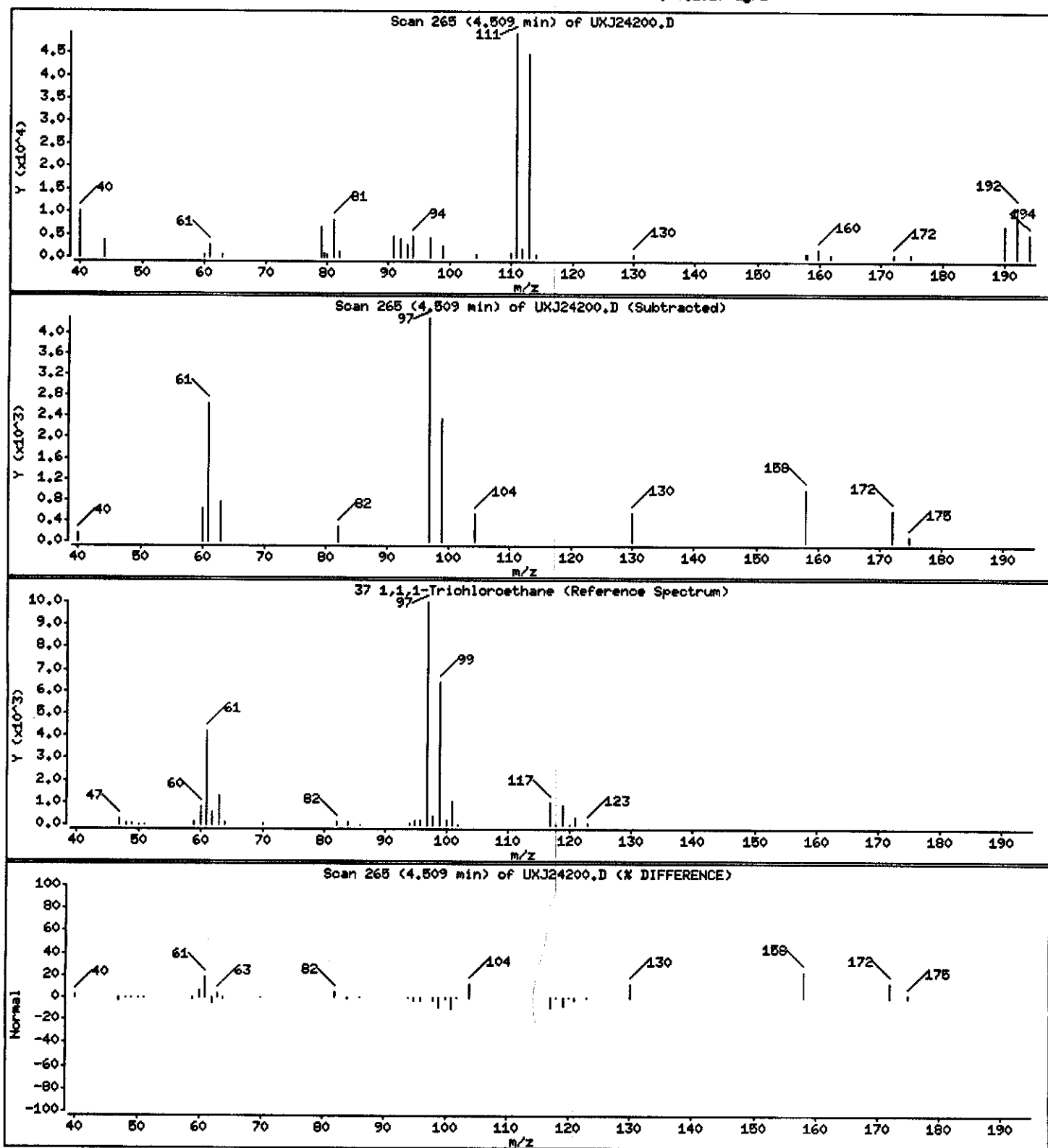
Operator: 43582

Column phase: DB624

Column diameter: 0.18

37 1,1,1-Trichloroethane

Concentration: 0.1927 ug/L



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J40929A.b\UXJ24200.D

Date : 29-SEP-2004 12:03

Client ID: VE539/12.5-17.5/092

Instrument: z3ux11.i

Sample Info: CQ67F1AA,5ML/5ML

Purge Volume: 5.0

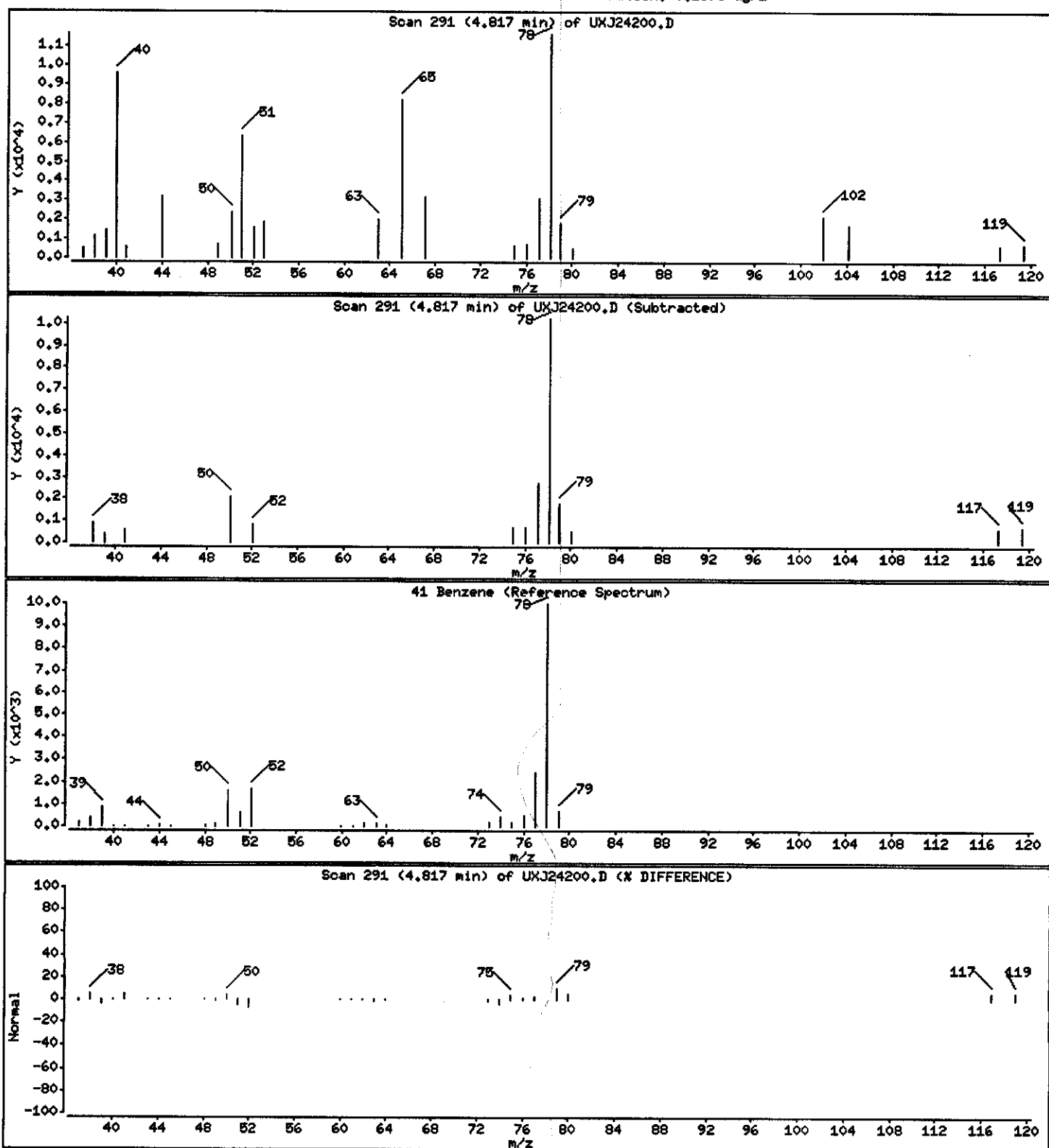
Operator: 43582

Column phase: DB624

Column diameter: 0.18

41 Benzene

Concentration: 0.1678 ug/L



Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J40929A.b\UXJ24200.D

Date : 29-SEP-2004 12:03

Client ID: VE539/12.5-17.5/092

Instrument: z3ux11.i

Sample Info: CQ67F1AA,5ML/5ML

Purge Volume: 5.0

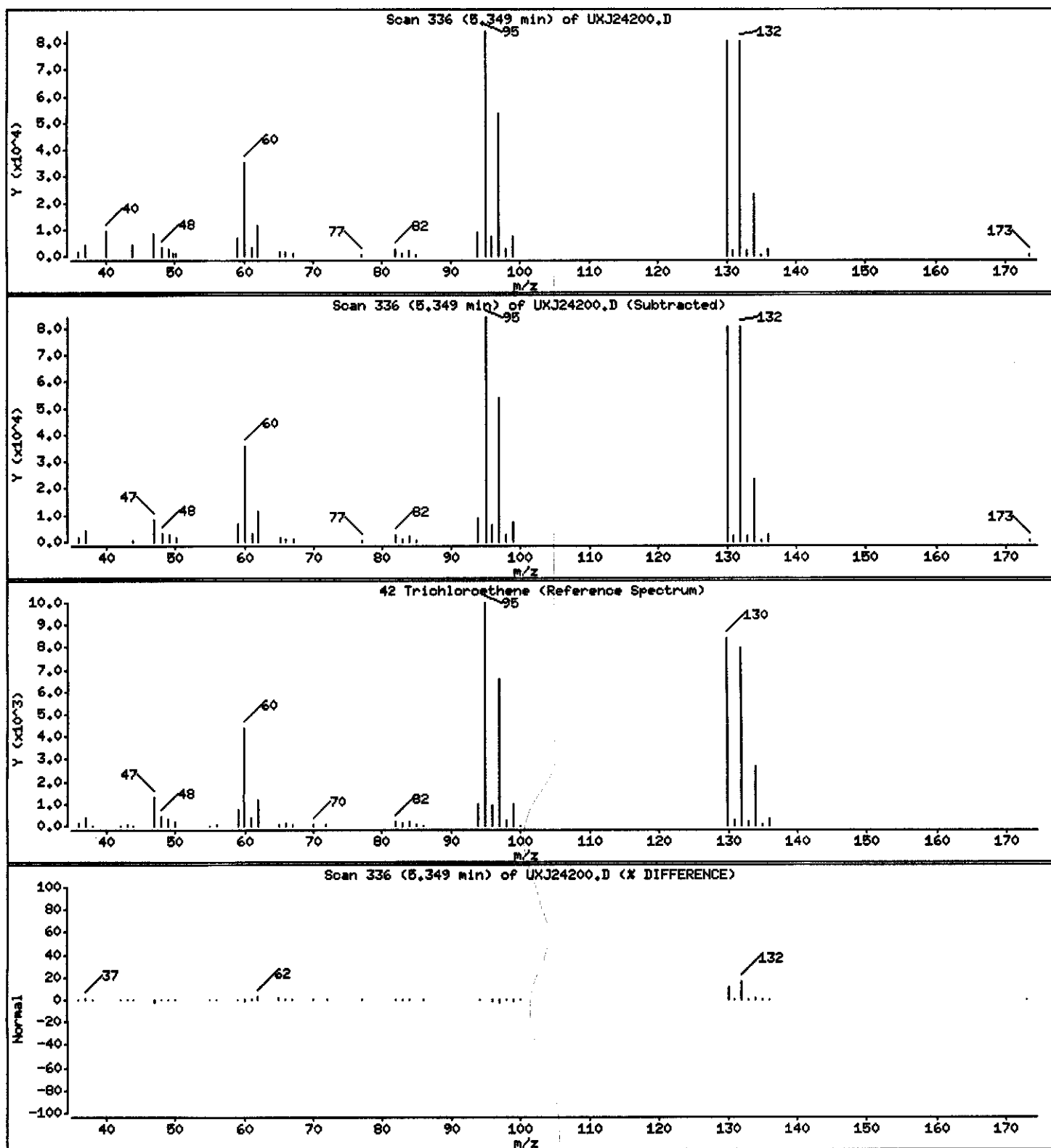
Operator: 43582

Column phase: DB624

Column diameter: 0.18

42 Trichloroethene

Concentration: 3.543 ug/L



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UXJ24200.D

Date : 29-SEP-2004 12:03

Client ID: VE539/12.5-17.5/092

Instrument: a3ux11.i

Sample Info: CQ67F1AA,5ML/5ML

Purge Volume: 5.0

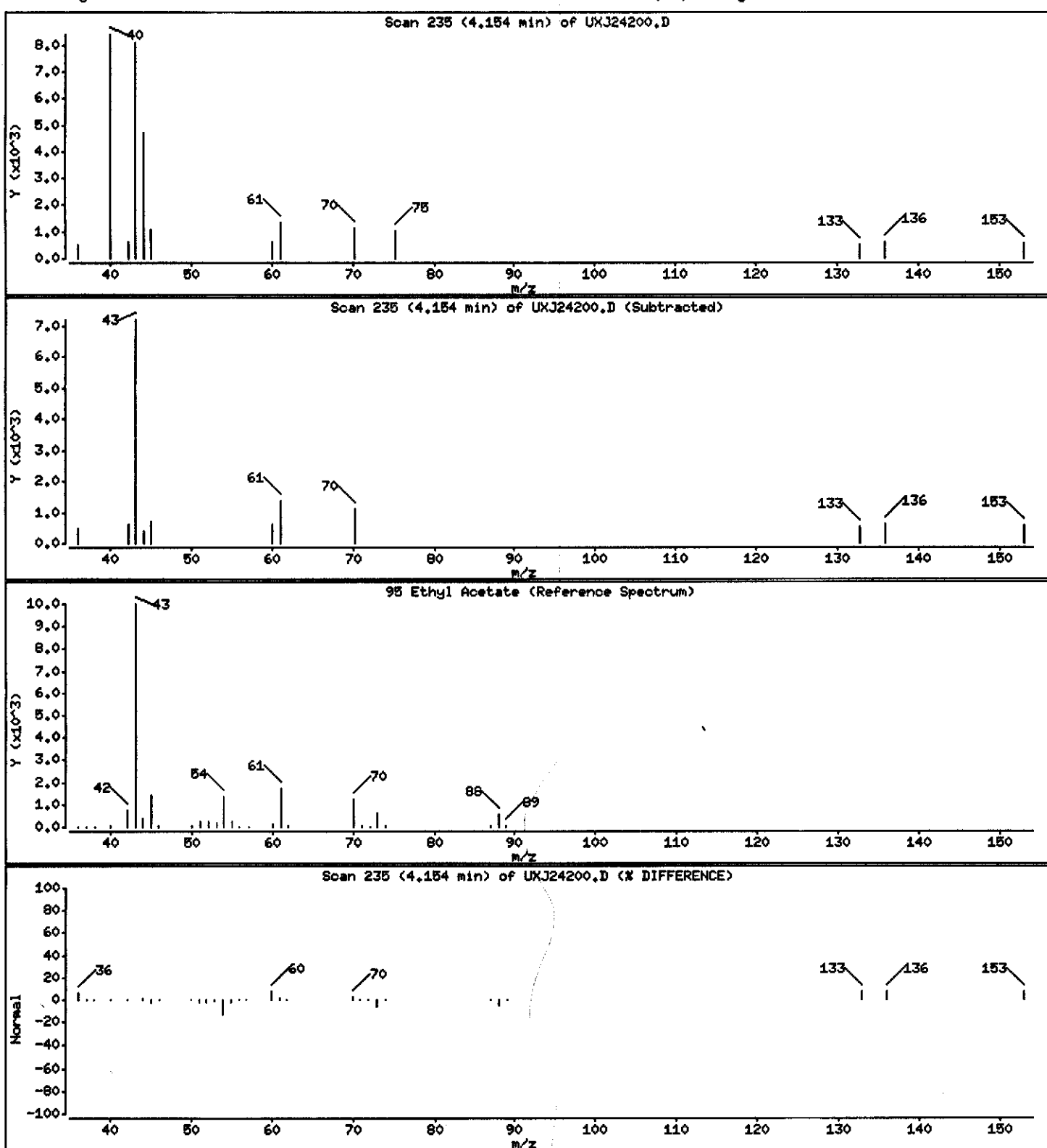
Operator: 43582

Column phase: DB624

Column diameter: 0.18

95 Ethyl Acetate

Concentration: 0.5832 ug/L



PAYNE FIRM INC.

Client Sample ID: TRIP BLANK/092404

GC/MS Volatiles

Lot-Sample #....: A4I270179-002 Work Order #....: GQ67G1AA Matrix.....: WQ
 Date Sampled....: 09/24/04 Date Received...: 09/25/04
 Prep Date.....: 09/29/04 Analysis Date...: 09/29/04
 Prep Batch #....: 4273268
 Dilution Factor: 1 Initial Wgt/Vol: 5 mL Final Wgt/Vol.: 5 mL
 Method.....: SW846 8260B

<u>PARAMETER</u>	<u>RESULT</u>	<u>REPORTING</u>	
		<u>LIMIT</u>	<u>UNITS</u>
Acetone	5.7 J	10	ug/L
Acetonitrile	ND	20	ug/L
Acrolein	ND	20	ug/L
Acrylonitrile	ND	20	ug/L
Benzene	ND	1.0	ug/L
Bromodichloromethane	ND	1.0	ug/L
Bromoform	ND	1.0	ug/L
Bromomethane	ND	1.0	ug/L
2-Butanone	2.5 J	10	ug/L
Carbon disulfide	ND	1.0	ug/L
Carbon tetrachloride	ND	1.0	ug/L
Chlorobenzene	ND	1.0	ug/L
Chloroprene	ND	2.0	ug/L
Dibromochloromethane	ND	1.0	ug/L
Chloroethane	ND	1.0	ug/L
Chloroform	ND	1.0	ug/L
Chloromethane	0.42 J,B	1.0	ug/L
3-Chloropropene	ND	2.0	ug/L
1,2-Dibromo-3-chloropropane (DBCP)	ND	2.0	ug/L
1,2-Dibromoethane	ND	1.0	ug/L
Dibromomethane	ND	1.0	ug/L
trans-1,4-Dichloro-2-butene	ND	1.0	ug/L
1,1-Dichloroethane	ND	1.0	ug/L
1,2-Dichloroethane	ND	1.0	ug/L
cis-1,2-Dichloroethene	ND	1.0	ug/L
trans-1,2-Dichloroethene	ND	1.0	ug/L
1,1-Dichloroethene	ND	1.0	ug/L
1,2-Dichloroethene (total)	ND	2.0	ug/L
Dichlorofluoromethane	ND	2.0	ug/L
1,2-Dichloropropane	ND	1.0	ug/L
cis-1,3-Dichloropropene	ND	1.0	ug/L
trans-1,3-Dichloropropene	ND	1.0	ug/L
1,4-Dioxane	ND	50	ug/L
Ethylbenzene	ND	1.0	ug/L
Ethyl methacrylate	ND	1.0	ug/L

(Continued on next page)

PAYNE FIRM INC.

Client Sample ID: TRIP BLANK/092404

GC/MS Volatiles

Lot-Sample #...: A4I270179-002 Work Order #...: GQ67G1AA Matrix.....: WQ

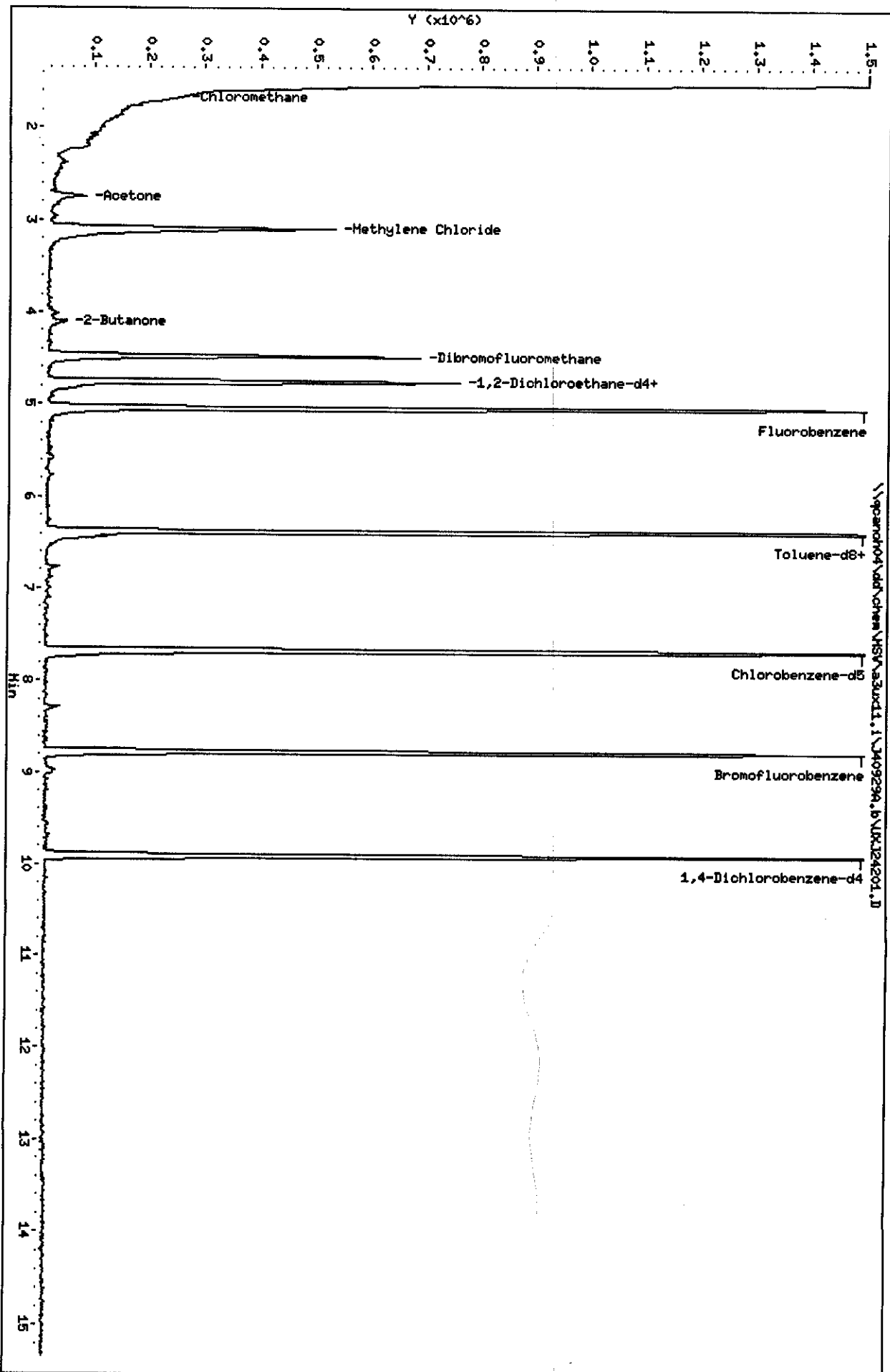
<u>PARAMETER</u>	<u>RESULT</u>	<u>REPORTING</u>	
		<u>LIMIT</u>	<u>UNITS</u>
2-Hexanone	ND	10	ug/L
Iodomethane	ND	1.0	ug/L
Isobutanol	ND	50	ug/L
Methacrylonitrile	ND	2.0	ug/L
Methylene chloride	7.2	1.0	ug/L
Methyl methacrylate	ND	2.0	ug/L
4-Methyl-2-pentanone	ND	10	ug/L
Propionitrile	ND	4.0	ug/L
Styrene	ND	1.0	ug/L
1,1,1,2-Tetrachloroethane	ND	1.0	ug/L
1,1,2,2-Tetrachloroethane	ND	1.0	ug/L
Tetrachloroethene	ND	1.0	ug/L
Toluene	0.19 J	1.0	ug/L
1,1,1-Trichloroethane	ND	1.0	ug/L
1,1,2-Trichloroethane	ND	1.0	ug/L
Trichloroethene	ND	1.0	ug/L
Trichlorofluoromethane	ND	1.0	ug/L
1,2,3-Trichloropropane	ND	1.0	ug/L
Vinyl acetate	ND	2.0	ug/L
Vinyl chloride	ND	1.0	ug/L
Xylenes (total)	ND	2.0	ug/L

<u>SURROGATE</u>	<u>PERCENT</u>	<u>RECOVERY</u>	
		<u>RECOVERY</u>	<u>LIMITS</u>
Dibromofluoromethane	118	(73 - 122)	
1,2-Dichloroethane-d4	113	(61 - 128)	
Toluene-d8	89	(76 - 110)	
4-Bromofluorobenzene	83	(74 - 116)	

NOTE(S) :

J Estimated result. Result is less than RL.

B Method blank contamination. The associated method blank contains the target analyte at a reportable level.



Data File: \\pcmanh04\dd\chem\MSA\z30x11.i\\J40929A.b\\J24201.D
 Date : 29-SEP-2004 12:26
 Client ID: TRIP BLANK\092404
 Sample Info: Q667G000,5ML5HL
 Purge Volume: 5.0
 Column Phase: DB624

 Instrument: z30x11.i
 Operator: 43582
 Column diameter: 0.18

 \\pcmanh04\dd\chem\MSA\z30x11.i\\J40929A.b\\J24201.D

Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40929A.b\UXJ24201.D
Report Date: 30-Sep-2004 08:44

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40929A.b\UXJ24201.D
Lab Smp Id: GQ67G1AA Client Smp ID: TRIP BLANK/092404
Inj Date : 29-SEP-2004 12:26
Operator : 43582 Inst ID: A3UX11.i
Smp Info : GQ67G1AA,5ML/5ML
Misc Info : J40929A,8260LLUX11,,43582
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 13
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
* 1 Fluorobenzene	96	5.041	5.041 (1.000)	1816483	50.0000		
* 2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1610955	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	682806	50.0000		
\$ 4 Dibromofluoromethane	113	4.485	4.473 (0.890)	500727	59.2062	11.841	
\$ 5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	654879	56.3119	11.262	
\$ 6 Toluene-d8	98	6.378	6.378 (0.831)	1724163	44.5502	8.910	
\$ 7 Bromofluorobenzene	95	8.780	8.780 (1.143)	684335	41.6162	8.323	
8 Dichlorodifluoromethane	85		Compound Not Detected.				
9 Chloromethane	50	1.669	1.692 (0.331)	35437	2.08997	0.4180	
10 Vinyl Chloride	62		Compound Not Detected.				
11 Bromomethane	94		Compound Not Detected.				
12 Chloroethane	64		Compound Not Detected.				
13 Trichlorofluoromethane	101		Compound Not Detected.				
15 Acrolein	56		Compound Not Detected.				
16 Acetone	43	2.745	2.734 (0.545)	131444	28.4839	5.697	
17 1,1-Dichloroethene	96		Compound Not Detected.				
18 Freon-113	151		Compound Not Detected.				

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24201.D
 Report Date: 30-Sep-2004 08:44

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
19 Iodomethane	142					Compound Not Detected.	
20 Carbon Disulfide	76					Compound Not Detected.	
21 Methylene Chloride	84		3.100	3.089 (0.615)		438175	35.9157 7.183
22 Acetonitrile	41					Compound Not Detected.	
23 Acrylonitrile	53					Compound Not Detected.	
24 Methyl tert-butyl ether	73					Compound Not Detected.	
25 trans-1,2-Dichloroethene	96					Compound Not Detected.	
26 Hexane	86					Compound Not Detected.	
27 Vinyl acetate	43					Compound Not Detected.	
28 1,1-Dichloroethane	63					Compound Not Detected.	
29 tert-Butyl Alcohol	59					Compound Not Detected.	
30 2-Butanone	43		4.106	4.094 (0.815)		61695	12.4002 2.480
M 31 1,2-Dichloroethene (total)	96					Compound Not Detected.	
32 cis-1,2-dichloroethene	96					Compound Not Detected.	
33 2,2-Dichloropropane	77					Compound Not Detected.	
34 Bromochloromethane	128					Compound Not Detected.	
35 Chloroform	83					Compound Not Detected.	
36 Tetrahydrofuran	42					Compound Not Detected.	
37 1,1,1-Trichloroethane	97					Compound Not Detected.	
38 1,1-Dichloropropene	75					Compound Not Detected.	
39 Carbon Tetrachloride	117					Compound Not Detected.	
40 1,2-Dichloroethane	62					Compound Not Detected.	
41 Benzene	78		4.816	4.816 (0.955)		34390	0.82231 0.1645
42 Trichloroethene	130					Compound Not Detected.	
43 1,2-Dichloropropane	63					Compound Not Detected.	
44 1,4-Dioxane	88					Compound Not Detected.	
45 Dibromomethane	93					Compound Not Detected.	
46 Bromodichloromethane	83					Compound Not Detected.	
47 2-Chloroethyl vinyl ether	63					Compound Not Detected.	
48 cis-1,3-Dichloropropene	75					Compound Not Detected.	
49 4-Methyl-2-pentanone	43					Compound Not Detected.	
50 Toluene	91		6.437	6.437 (0.838)		46118	0.95856 0.1917
51 trans-1,3-Dichloropropene	75					Compound Not Detected.	
52 Ethyl Methacrylate	69					Compound Not Detected.	
53 1,1,2-Trichloroethane	97					Compound Not Detected.	
54 1,3-Dichloropropane	76					Compound Not Detected.	
55 Tetrachloroethene	164					Compound Not Detected.	
56 2-Hexanone	43					Compound Not Detected.	
57 Dibromochloromethane	129					Compound Not Detected.	
58 1,2-Dibromoethane	107					Compound Not Detected.	
59 Chlorobenzene	112					Compound Not Detected.	
60 1,1,1,2-Tetrachloroethane	131					Compound Not Detected.	
61 Ethylbenzene	106					Compound Not Detected.	
62 m + p-Xylene	106					Compound Not Detected.	
M 63 Xylenes (total)	106					Compound Not Detected.	
64 Xylene-o	106					Compound Not Detected.	
65 Styrene	104					Compound Not Detected.	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24201.D
 Report Date: 30-Sep-2004 08:44

Compounds	QUANT SIG	MASS	CONCENTRATIONS				
			RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
66 Bromoform	---	173	---	---	---	Compound Not Detected.	-----
67 Isopropylbenzene	---	105	---	---	---	Compound Not Detected.	-----
68 1,1,2,2-Tetrachloroethane	---	83	---	---	---	Compound Not Detected.	-----
69 1,4-Dichloro-2-butene	---	53	---	---	---	Compound Not Detected.	-----
70 1,2,3-Trichloropropane	---	110	---	---	---	Compound Not Detected.	-----
71 Bromobenzene	---	156	---	---	---	Compound Not Detected.	-----
72 n-Propylbenzene	---	120	---	---	---	Compound Not Detected.	-----
73 2-Chlorotoluene	---	126	---	---	---	Compound Not Detected.	-----
74 1,3,5-Trimethylbenzene	---	105	---	---	---	Compound Not Detected.	-----
75 4-Chlorotoluene	---	126	---	---	---	Compound Not Detected.	-----
76 tert-Butylbenzene	---	119	---	---	---	Compound Not Detected.	-----
77 1,2,4-Trimethylbenzene	---	105	---	---	---	Compound Not Detected.	-----
78 sec-Butylbenzene	---	105	---	---	---	Compound Not Detected.	-----
79 4-Isopropyltoluene	---	119	---	---	---	Compound Not Detected.	-----
80 1,3-Dichlorobenzene	---	146	---	---	---	Compound Not Detected.	-----
81 1,4-Dichlorobenzene	---	146	---	---	---	Compound Not Detected.	-----
82 n-Butylbenzene	---	91	---	---	---	Compound Not Detected.	-----
83 1,2-Dichlorobenzene	---	146	---	---	---	Compound Not Detected.	-----
84 1,2-Dibromo-3-chloropropane	---	157	---	---	---	Compound Not Detected.	-----
85 1,2,4-Trichlorobenzene	---	180	---	---	---	Compound Not Detected.	-----
86 Hexachlorobutadiene	---	225	---	---	---	Compound Not Detected.	-----
87 Naphthalene	---	128	---	---	---	Compound Not Detected.	-----
88 1,2,3-Trichlorobenzene	---	180	---	---	---	Compound Not Detected.	-----
14 Dichlorofluoromethane	---	67	---	---	---	Compound Not Detected.	-----
89 Ethyl Ether	---	59	---	---	---	Compound Not Detected.	-----
91 3-Chloropropene	---	76	---	---	---	Compound Not Detected.	-----
92 Isopropyl Ether	---	87	---	---	---	Compound Not Detected.	-----
93 2-Chloro-1,3-butadiene	---	53	---	---	---	Compound Not Detected.	-----
94 Propionitrile	---	54	---	---	---	Compound Not Detected.	-----
95 Ethyl Acetate	---	43	---	---	---	Compound Not Detected.	-----
96 Methacrylonitrile	---	41	---	---	---	Compound Not Detected.	-----
97 Isobutanol	---	41	---	---	---	Compound Not Detected.	-----
99 n-Butanol	---	56	---	---	---	Compound Not Detected.	-----
100 Methyl Methacrylate	---	41	---	---	---	Compound Not Detected.	-----
101 2-Nitropropane	---	41	---	---	---	Compound Not Detected.	-----
103 Cyclohexanone	---	55	---	---	---	Compound Not Detected.	-----
98 Cyclohexane	---	56	---	---	---	Compound Not Detected.	-----
143 Methyl Acetate	---	43	---	---	---	Compound Not Detected.	-----
144 Methylcyclohexane	---	83	---	---	---	Compound Not Detected.	-----
141 1,3,5-Trichlorobenzene	---	180	---	---	---	Compound Not Detected.	-----

Data File: \\qcanoh04\dd\chem\MSV\z3ux11.i\J40929A.b\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: z3ux11.i

Sample Info: CQ67C1AA,5ML/5ML

Purge Volume: 5.0

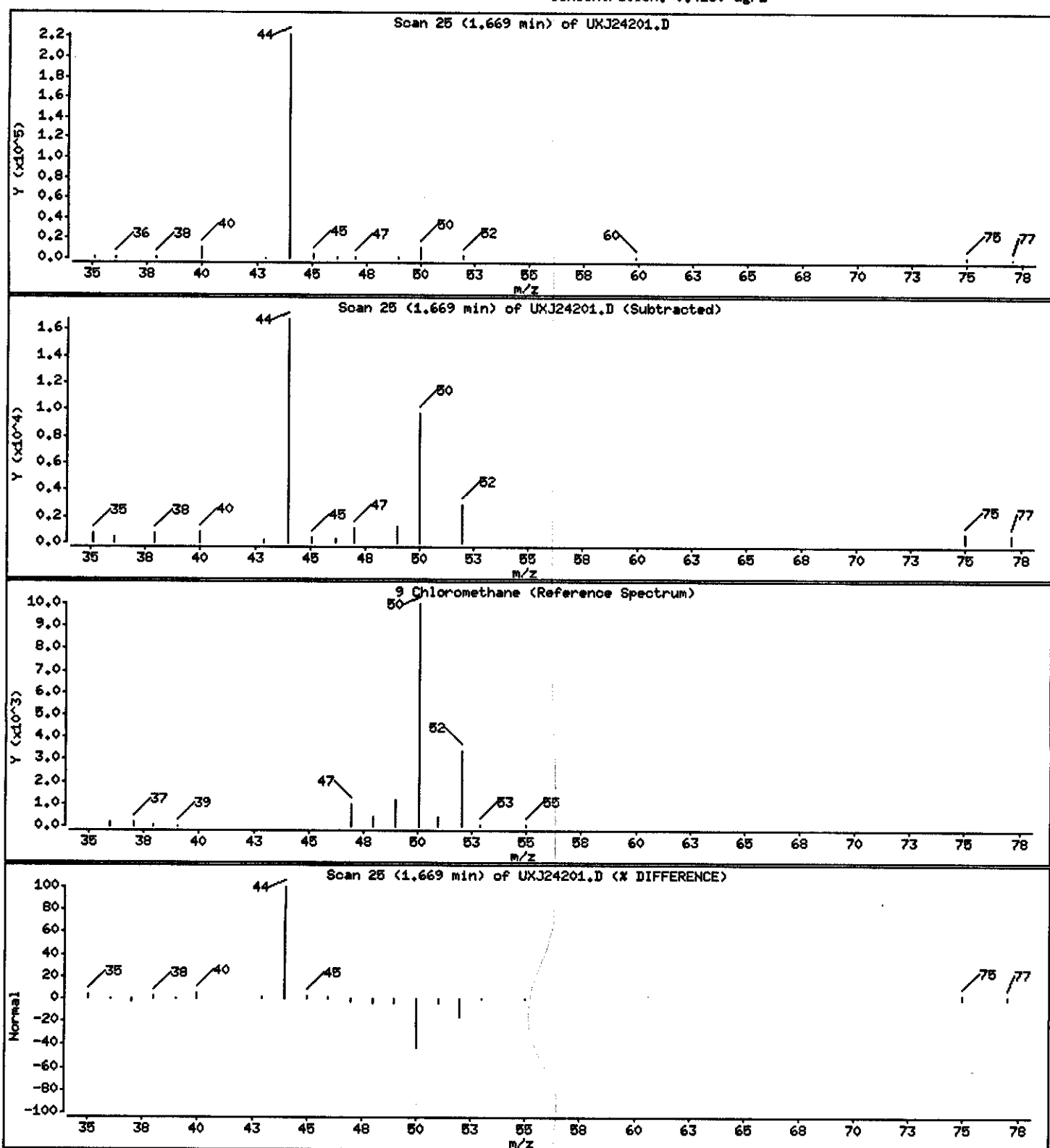
Operator: 43582

Column phase: DB624

Column diameter: 0.18

9 Chloromethane

Concentration: 0.4180 ug/L



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: a3ux11.i

Sample Info: CQ67G1AA,5ML/5ML

Purge Volume: 5.0

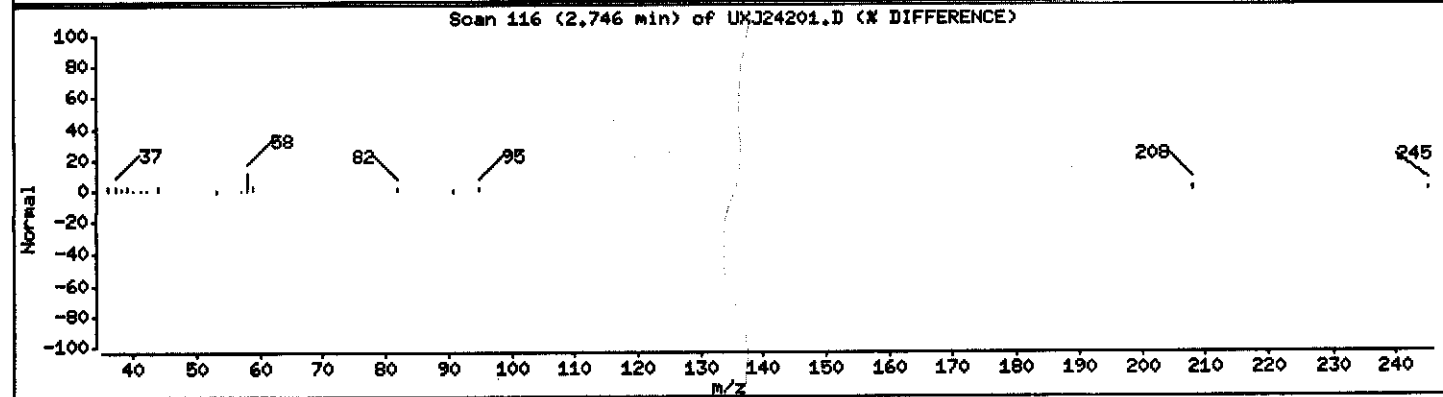
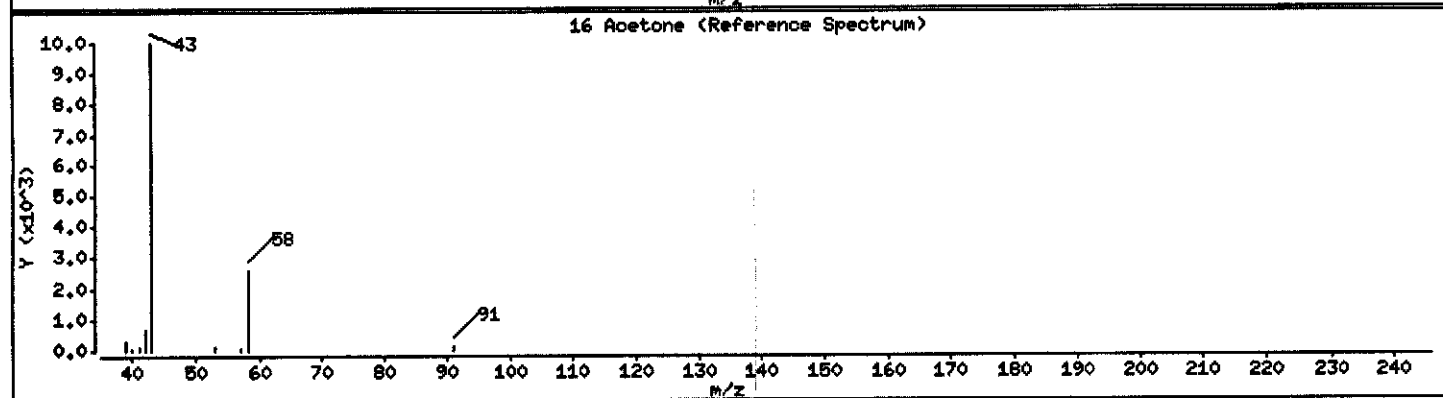
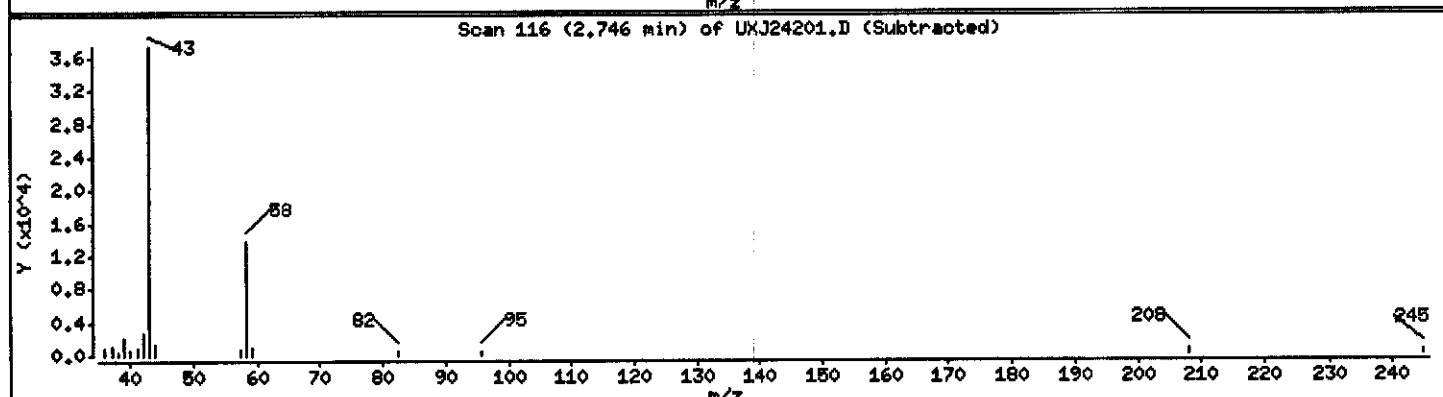
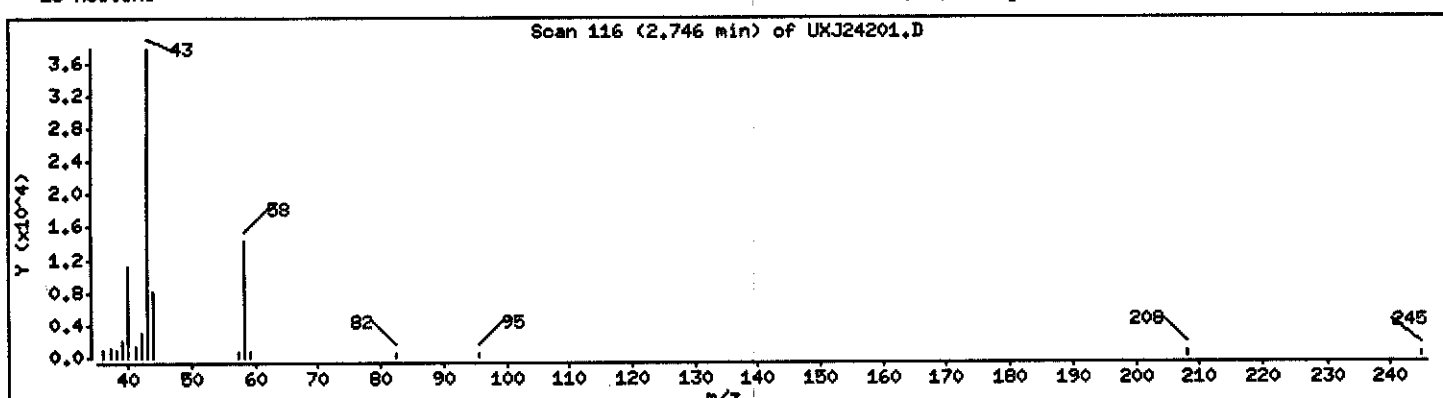
Operator: 43582

Column phase: DB624

Column diameter: 0.18

16 Acetone

Concentration: 5.697 ug/L



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: a3ux11.i

Sample Info: CQ67G1AA,5ML/5ML

Operator: 43582

Purge Volume: 5.0

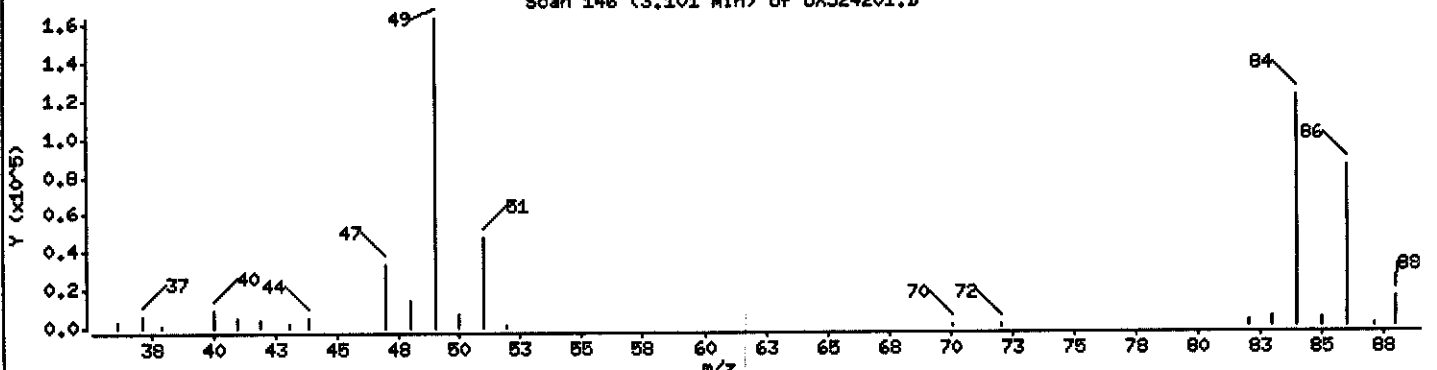
Column diameter: 0.18

Column phase: DB624

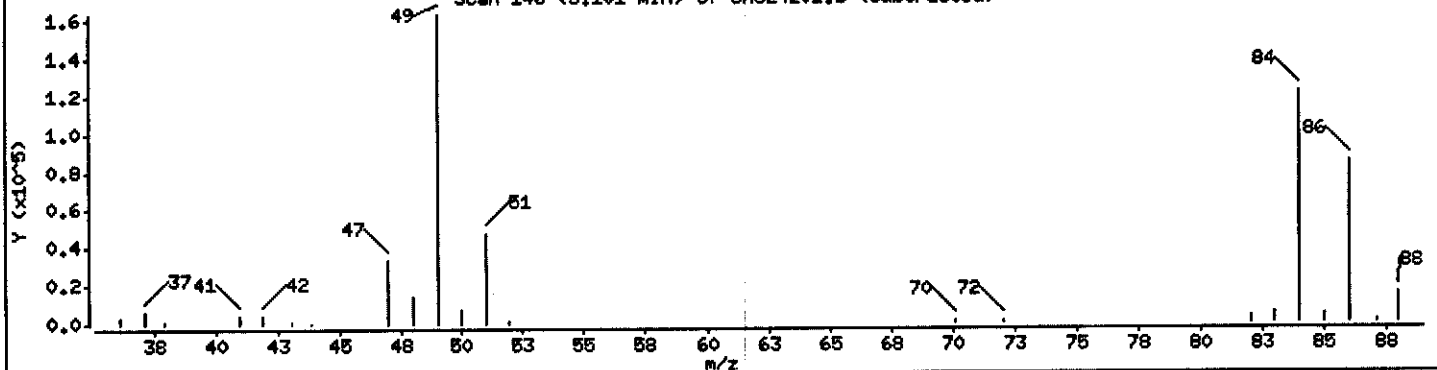
Concentration: 7.183 ug/L

21. Methylene Chloride

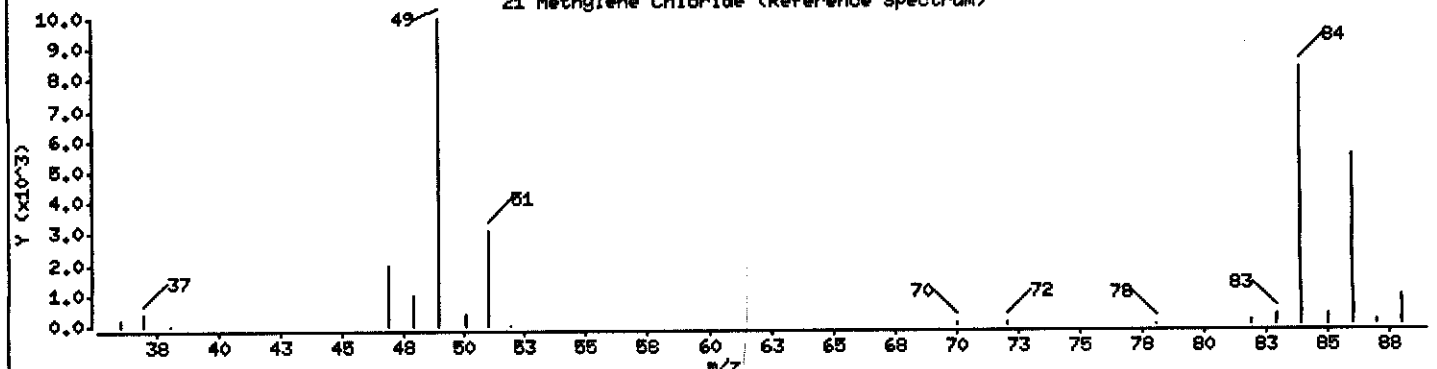
Scan 146 (3.101 min) of UXJ24201.D



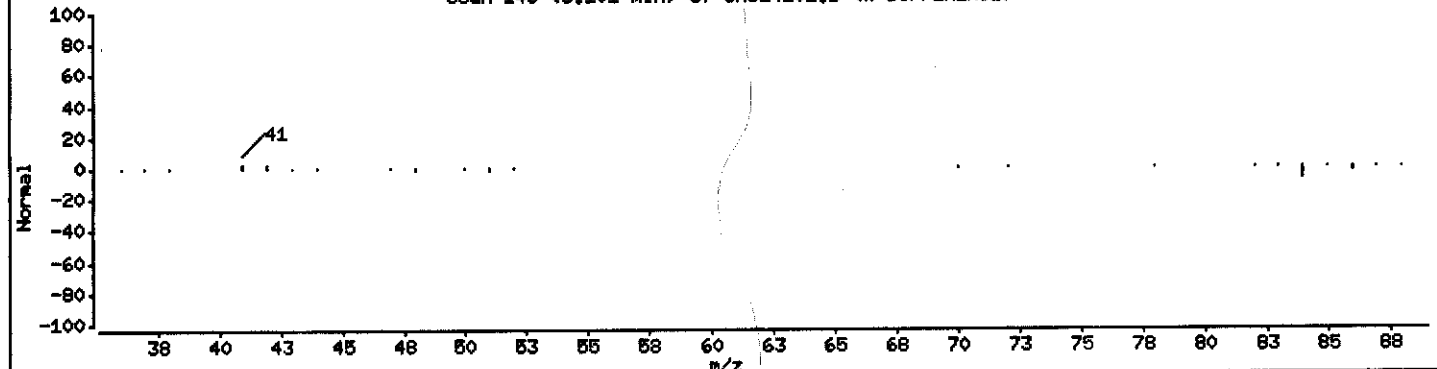
Scan 146 (3.101 min) of UXJ24201.D (Subtracted)



21 Methylene Chloride (Reference Spectrum)



Scan 146 (3.101 min) of UXJ24201.D (% DIFFERENCE)



Data File: \\qcanch04\dd\chem\MSV\z3ux11.i\J40929A.b\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: z3ux11.i

Sample Info: GQ67C1AA,5ML/5ML

Purge Volume: 5.0

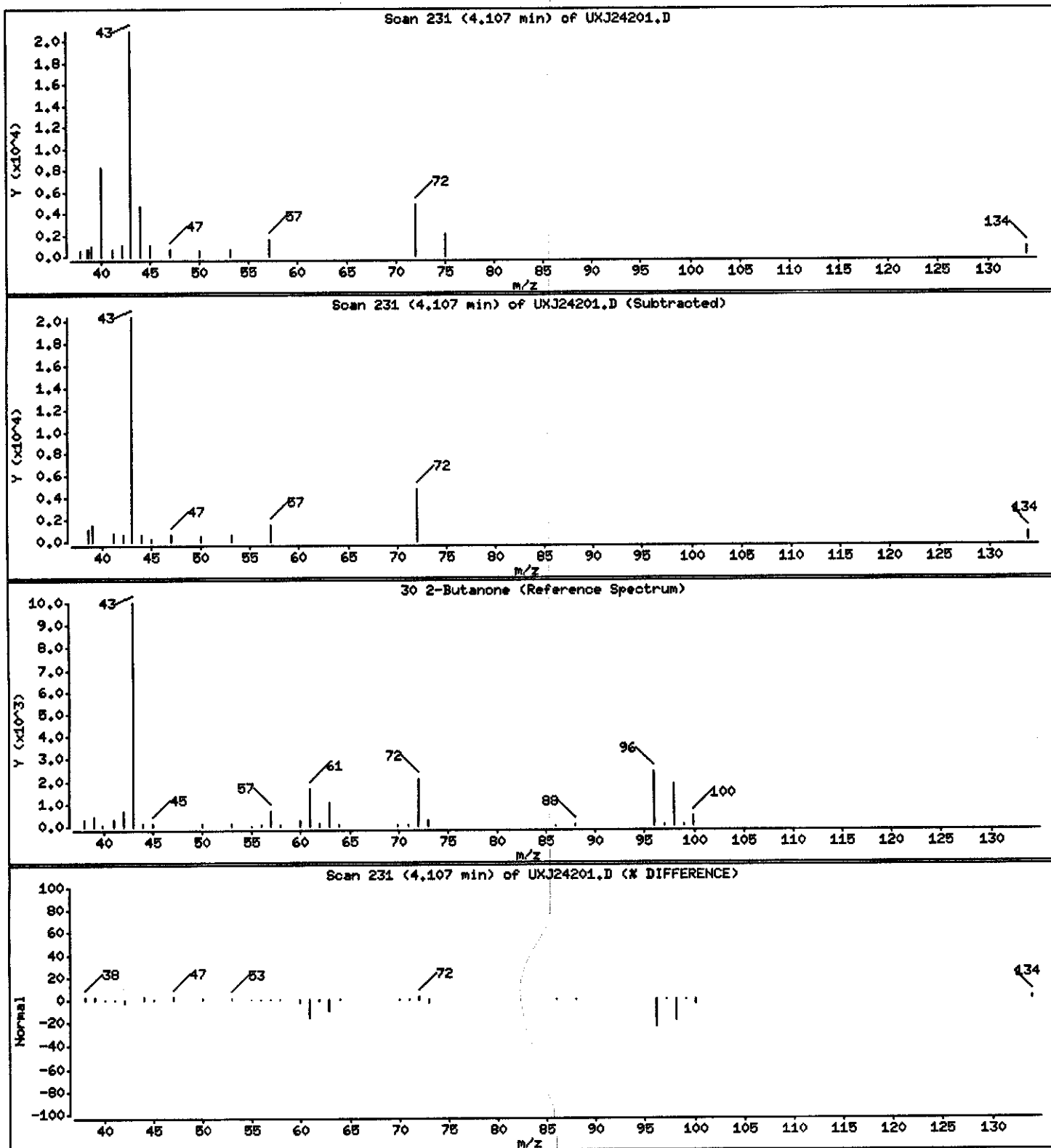
Operator: 43582

Column phaset: DB624

Column diameter: 0.18

30 2-Butanone

Concentration: 2.480 ug/L



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: a3ux11.i

Sample Info: CQ67G1AA,5ML/5ML

Purge Volume: 5.0

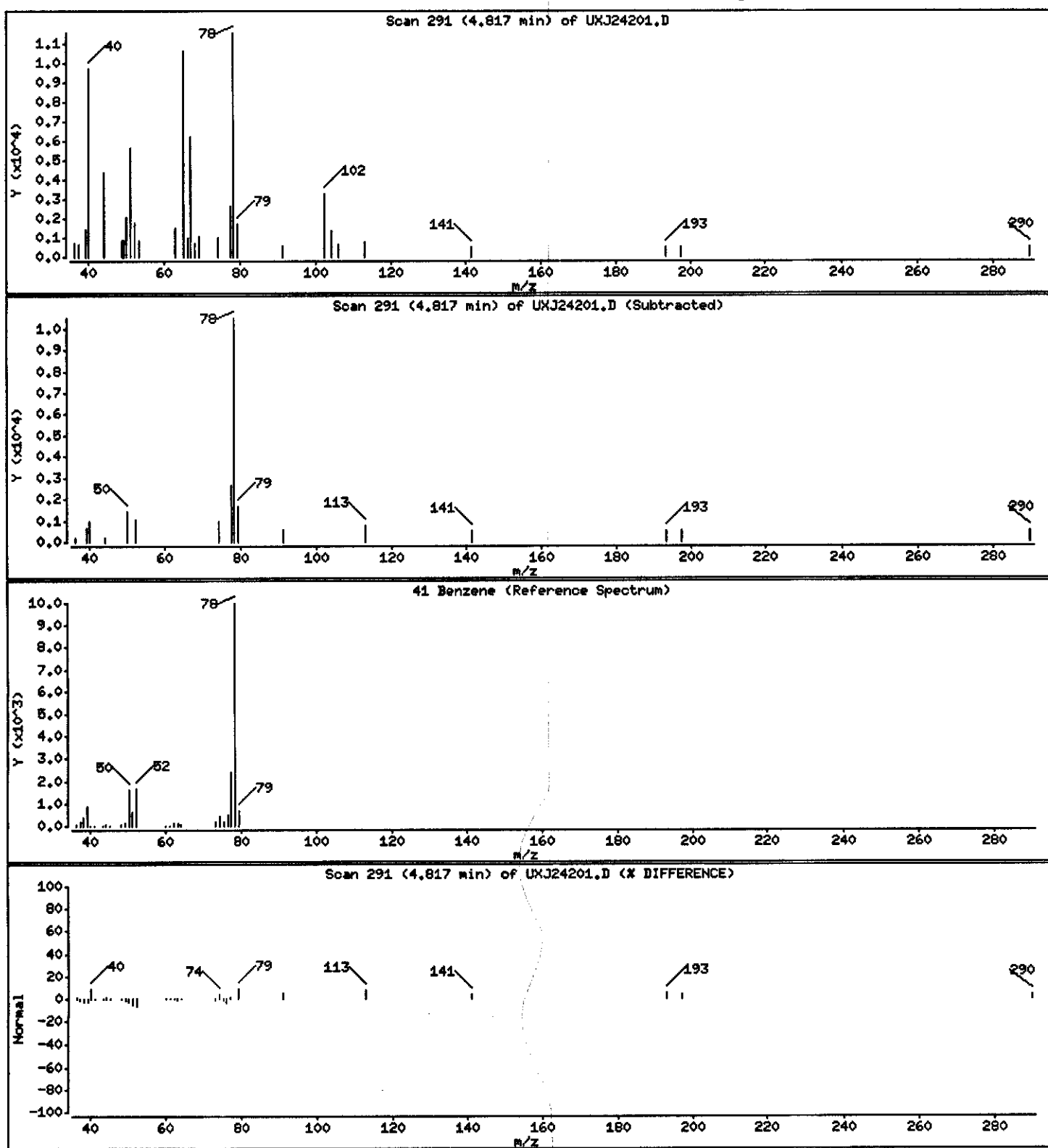
Operator: 43582

Column phase: DB624

Column diameter: 0.18

41 Benzene

Concentration: 0.1645 ug/L



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J40929A.b\UXJ24201.D

Date : 29-SEP-2004 12:26

Client ID: TRIP BLANK/092404

Instrument: s3ux11.i

Sample Info: CQ67C1AA,5ML/5ML

Purge Volume: 5.0

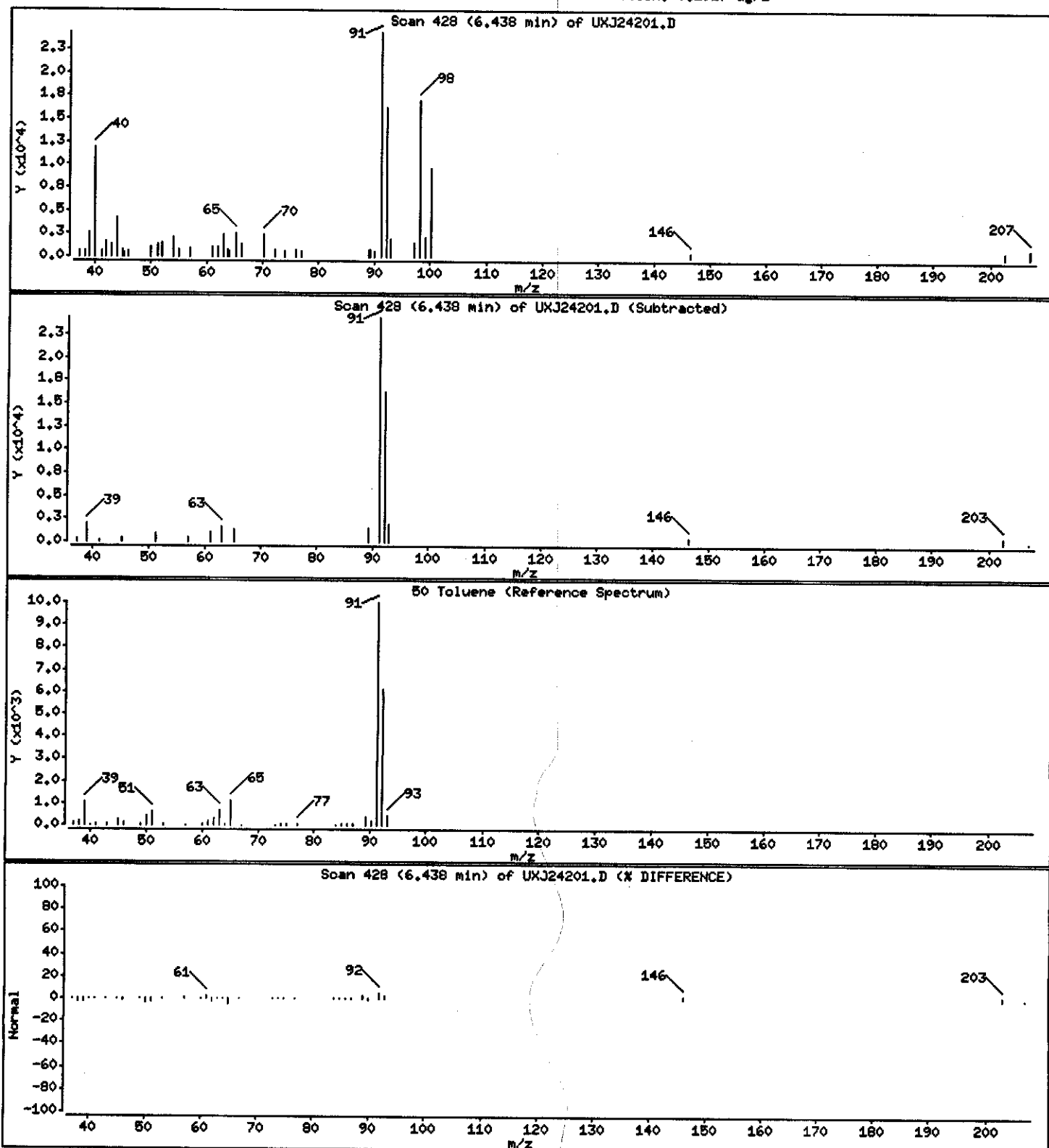
Operator: 43582

Column phase: DB624

Column diameter: 0.18

50 Toluene

Concentration: 0.1917 ug/L



STANDARD DATA

Report Date: 11-19-04

Calibration History

Method : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Start Cal Date: 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Last Cal Level: 1
Last Cal Type : Initial Calibration

Initial Calibration

Injection Date	Sublist	Calibration File
Cal Level: 1 , Cal Amount: 5.000		
14-SEP-2004 15:41	2-8260	UXJ23875.D
16-AUG-2004 18:11	3-IX	UXJ23214.D
Cal Level: 2 , Cal Amount: 10.000		
14-SEP-2004 15:19	2-8260	UXJ23874.D
16-AUG-2004 17:48	3-IX	UXJ23213.D
Cal Level: 3 , Cal Amount: 25.000		
14-SEP-2004 14:57	2-8260	UXJ23873.D
16-AUG-2004 17:26	3-IX	UXJ23212.D
Cal Level: 4 , Cal Amount: 50.000		
14-SEP-2004 14:33	2-8260	UXJ23872.D
16-AUG-2004 17:03	3-IX	UXJ23211.D
Cal Level: 5 , Cal Amount: 100.00		
14-SEP-2004 14:10	2-8260	UXJ23871.D
16-AUG-2004 16:40	3-IX	UXJ23210.D
Cal Level: 6 , Cal Amount: 200.00		
14-SEP-2004 13:48	2-8260	UXJ23870.D
16-AUG-2004 16:18	3-IX	UXJ23209.D

Continuing Calibration

14-SEP-2004 14:33	2-8260	UXJ23872.D

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

Calibration File Names:

Level 1: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23875.D
 Level 2: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23874.D
 Level 3: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23873.D
 Level 4: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23872.D
 Level 5: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23871.D
 Level 6: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23870.D

Compound	5.000	10.000	25.000	50.000	100.000	200.000			
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	% RSD	
8 Dichlorodifluoromethane	0.28836	0.23754	0.28173	0.25161	0.24515	0.26401	0.26140	7.796	
9 Chloromethane	0.55674	0.49898	0.45483	0.43875	0.42154	0.42948	0.46672	11.132	
10 Vinyl Chloride	0.32349	0.31847	0.33246	0.31180	0.29804	0.33125	0.31925	4.066	
11 Bromomethane	0.17259	0.16446	0.14190	0.15123	0.13819	0.14020	0.15143	9.386	
12 Chloroethane	0.23244	0.24693	0.24362	0.23419	0.22089	0.22969	0.23463	4.043	
13 Trichlorofluoromethane	0.38407	0.31147	0.34717	0.32533	0.30602	0.33365	0.33462	8.499	
14 Dichlorofluoromethane	0.45871	0.50000	0.48561	0.49154	0.48286	0.49097	0.48495	2.915	
15 Acrolein	0.03199	0.03065	0.03129	0.03093	0.03156	0.03098	0.03123	1.551	
16 Acetone	0.16115	0.13958	0.11422	0.11384	0.10613	0.10203	0.12282	18.621	
17 1,1-Dichloroethene	0.24613	0.23009	0.19938	0.23265	0.20714	0.22623	0.22360	7.735	
18 Freon-113	0.16870	0.16626	0.10968	0.16762	0.13496	0.15980	0.15117	15.834	
19 Iodomethane	0.31149	0.34851	0.32631	0.33364	0.33020	0.33284	0.33050	3.627	
20 Carbon Disulfide	0.91244	0.87319	0.74270	0.85398	0.78522	0.84395	0.83525	7.360	
21 Methylene Chloride	0.75227	0.54750	0.35921	0.31815	0.29068	0.28051	0.42472	44.272	
22 Acetonitrile	0.03290	0.03019	0.03062	0.02865	0.02976	0.02592	0.02967	7.789	
23 Acrylonitrile	0.09137	0.09573	0.09318	0.09088	0.09229	0.09022	0.09228	2.153	
24 Methyl tert-butyl ether	0.65699	0.67280	0.74332	0.74432	0.74979	0.73394	0.71686	5.703	
25 trans-1,2-Dichloroethene	0.28262	0.27489	0.25328	0.26749	0.25617	0.25710	0.26526	4.435	
26 Hexane	0.05179	0.04861	0.03324	0.04941	0.04250	0.04921	0.04579	15.039	
27 Vinyl acetate	0.41519	0.38277	0.42613	0.42027	0.44948	0.45290	0.42446	6.033	
28 1,1-Dichloroethane	0.51364	0.49348	0.46074	0.48352	0.47362	0.47833	0.48389	3.753	
29 tert-Butyl Alcohol	0.02052	0.02018	0.01934	0.01924	0.01980	0.01761	0.01945	5.262	
30 2-Butanone	0.15894	0.12780	0.13677	0.13376	0.13540	0.12902	0.13695	8.281	
M 31 1,2-Dichloroethene (total)	0.28520	0.28491	0.26194	0.27101	0.26404	0.26479	0.27198	3.886	
32 cis-1,2-dichloroethene	0.28778	0.29492	0.27061	0.27453	0.27191	0.27248	0.27870	3.636	

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

Compound	5.000	10.000	25.000	50.000	100.000	200.000	RRF	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6		
33 2,2-Dichloropropane	0.29593	0.30774	0.27673	0.29383	0.28018	0.29523	0.29161	3.907
34 Bromochloromethane	0.12769	0.14200	0.12631	0.12865	0.12580	0.12388	0.12906	5.074
35 Chloroform	0.51012	0.50575	0.47005	0.48650	0.47366	0.47250	0.48643	3.631
36 Tetrahydrofuran	0.10018	0.08347	0.06844	0.07301	0.07510	0.07044	0.07844	15.112
37 1,1,1-Trichloroethane	0.39287	0.37451	0.33541	0.38300	0.34846	0.36998	0.36737	5.878
38 1,1-Dichloropropene	0.34921	0.35729	0.30656	0.35792	0.32912	0.35428	0.34240	6.001
39 Carbon Tetrachloride	0.30286	0.29259	0.24537	0.31080	0.27649	0.30261	0.28845	8.383
40 1,2-Dichloroethane	0.40025	0.42402	0.38489	0.39237	0.39169	0.39145	0.39745	3.498
41 Benzene	1.25995	1.19011	1.12471	1.13415	1.09548	1.10254	1.15116	5.467
42 Trichloroethene	0.27283	0.26390	0.25129	0.26732	0.25842	0.26258	0.26272	2.818
43 1,2-Dichloropropane	0.29669	0.30122	0.27194	0.28098	0.27691	0.27607	0.28397	4.241
44 1,4-Dioxane	0.00242	0.00233	0.00258	0.00261	0.00260	0.00206	0.00243	8.907
45 Dibromomethane	0.15285	0.16666	0.15233	0.15822	0.15862	0.15545	0.15736	3.340
46 Bromodichloromethane	0.38193	0.38487	0.36100	0.37042	0.36931	0.37381	0.37355	2.345
47 2-Chloroethyl vinyl ether	0.13950	0.14798	0.15985	0.16693	0.17284	0.17092	0.15967	8.385
48 cis-1,3-Dichloropropene	0.43417	0.47734	0.43633	0.45487	0.46634	0.47331	0.45706	4.058
49 4-Methyl-2-pentanone	0.23661	0.23635	0.24230	0.25213	0.25538	0.25182	0.24577	3.424
50 Toluene	1.48717	1.54897	1.47049	1.49192	1.46610	1.49497	1.49327	1.987
51 trans-1,3-Dichloropropene	0.50647	0.53118	0.51429	0.54236	0.55606	0.56198	0.53539	4.162
52 Ethyl Methacrylate	0.38934	0.45025	0.45000	0.48061	0.49453	0.49921	0.46066	8.865
53 1,1,2-Trichloroethane	0.30188	0.31382	0.29748	0.30352	0.30279	0.29789	0.30290	1.953
54 1,3-Dichloropropane	0.57206	0.59182	0.56392	0.55975	0.57182	0.56489	0.57071	1.996
55 Tetrachloroethene	0.25509	0.26206	0.22142	0.24509	0.22718	0.24073	0.24193	6.478
56 2-Hexanone	0.22234	0.23807	0.24231	0.24889	0.24606	0.25308	0.24179	4.486
57 Dibromochloromethane	0.30967	0.33372	0.31997	0.31951	0.32624	0.32930	0.32307	2.641
58 1,2-Dibromoethane	0.26921	0.31637	0.29041	0.30230	0.30732	0.30501	0.29844	5.561
59 Chlorobenzene	0.96525	1.03980	0.94396	0.94706	0.94928	0.95592	0.96688	3.777
60 1,1,1,2-Tetrachloroethane	0.33779	0.34414	0.32638	0.32925	0.34031	0.33757	0.33590	2.013
61 Ethylbenzene	0.46024	0.50768	0.46972	0.48906	0.48935	0.51358	0.48827	4.245
62 m + p-Xylene	0.60709	0.64675	0.61837	0.63926	0.62027	0.64155	0.62888	2.511
M 63 Xylenes (total)	0.59346	0.64358	0.61984	0.63276	0.62064	0.64001	0.62505	2.925
64 Xylene-o	0.56621	0.63725	0.62277	0.61975	0.62137	0.63694	0.61738	4.252
65 Styrene	1.00309	1.11779	1.08299	1.11462	1.13838	1.16736	1.10404	5.143

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
 End Cal Date : 14-SEP-2004 15:41
 Quant Method : ISTD
 Origin : Disabled
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\qcanoh04\dd\chem\MSV\aux11.i\J40914B-IC.b\8260LLUX11.m
 Cal Date : 14-Sep-2004 16:57 tapsvc
 Curve Type : Average

Compound	5.000	10.000	25.000	50.000	100.000	200.000	—	—	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	—	—
66 Bromoform	0.20132	0.22491	0.21511	0.21840	0.22531	0.23254	0.21960	4.920	
67 Isopropylbenzene	1.34498	1.36235	1.30493	1.41191	1.35691	1.46909	1.37503	4.180	
68 1,1,2,2-Tetrachloroethane	0.74074	0.78253	0.73952	0.75703	0.76116	0.74128	0.75371	2.237	
69 1,4-Dichloro-2-butene	0.23161	0.23946	0.23911	0.25172	0.26183	0.26419	0.24799	5.373	
70 1,2,3-Trichloropropane	0.25299	0.25205	0.24157	0.24830	0.24773	0.24135	0.24733	2.016	
71 Bromobenzene	0.74497	0.77908	0.72155	0.75186	0.74513	0.73336	0.74599	2.605	
72 n-Propylbenzene	0.71725	0.70753	0.62905	0.72290	0.67589	0.72046	0.69551	5.299	
73 2-Chlorotoluene	0.71622	0.69234	0.65887	0.70074	0.67558	0.68895	0.68879	2.887	
74 1,3,5-Trimethylbenzene	2.25968	2.26774	2.20534	2.39411	2.31559	2.44388	2.31439	3.874	
75 4-Chlorotoluene	0.76654	0.75820	0.70598	0.74482	0.71695	0.72231	0.73580	3.304	
76 tert-Butylbenzene	1.89049	1.94223	1.70074	1.94123	1.83516	1.94010	1.87499	5.079	
77 1,2,4-Trimethylbenzene	2.33163	2.49724	2.33803	2.57507	2.50096	2.58086	2.47063	4.492	
78 sec-Butylbenzene	2.66025	2.55624	2.28546	2.65769	2.44749	2.62040	2.53792	5.801	
79 4-Isopropyltoluene	2.02816	2.10772	1.90515	2.27140	2.10588	2.25071	2.11151	6.511	
80 1,3-Dichlorobenzene	1.50394	1.41628	1.30596	1.36727	1.33637	1.33353	1.37723	5.270	
81 1,4-Dichlorobenzene	1.49216	1.50657	1.37893	1.45011	1.40876	1.40879	1.44089	3.529	
82 n-Butylbenzene	1.86201	1.92154	1.64744	2.01989	1.84399	2.02271	1.88626	7.389	
83 1,2-Dichlorobenzene	1.44948	1.40634	1.30068	1.35160	1.32364	1.30712	1.35648	4.397	
84 1,2-Dibromo-3-chloropropane	0.12641	0.12593	0.12612	0.12831	0.13070	0.12832	0.12763	1.446	
85 1,2,4-Trichlorobenzene	0.56391	0.60976	0.52246	0.58135	0.57921	0.60830	0.57750	5.596	
86 Hexachlorobutadiene	0.39076	0.31635	0.25219	0.27519	0.23759	0.23788	0.28500	20.941	
87 Naphthalene	1.19572	1.26888	1.21865	1.39469	1.53842	1.56161	1.36300	11.780	
88 1,2,3-Trichlorobenzene	0.38318	0.44192	0.36462	0.41931	0.44173	0.43401	0.41413	7.910	
89 Ethyl Ether	0.26862	0.25572	0.24190	0.23691	0.23958	0.23653	0.24654	5.241	
90 Ethanol	+++++	+++++	+++++	+++++	+++++	+++++	+++++	++	<-
91 3-Chloropropene	0.08501	0.09752	0.10420	0.10696	0.10892	0.11570	0.10305	10.335	
92 Isopropyl Ether	0.19894	0.21144	0.22424	0.23464	0.23633	0.23561	0.22353	6.886	
93 2-Chloro-1,3-butadiene	0.33010	0.35251	0.38261	0.39081	0.38671	0.39380	0.37276	6.882	
94 Propionitrile	0.04383	0.04396	0.04410	0.04111	0.04094	0.03990	0.04231	4.404	
95 Ethyl Acetate	0.25153	0.23918	0.24678	0.23828	0.24450	0.25022	0.24508	2.253	
96 Methacrylonitrile	0.15470	0.15983	0.15530	0.15862	0.16298	0.16195	0.15890	2.134	
97 Isobutanol	0.01214	0.01067	0.01151	0.01109	0.01159	0.01152	0.01142	4.362	
98 Cyclohexane	0.37486	0.38746	0.29089	0.40866	0.34684	0.39833	0.36784	11.783	

Report Date : 14-Sep-2004 17:02

STL North Canton

INITIAL CALIBRATION DATA

Start Cal Date : 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Quant Method : ISTD
Origin : Disabled
Target Version : 4.04
Integrator : HP RTE
Method file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\8260LLUX11.m
Cal Date : 14-Sep-2004 16:57 tapsvc
Curve Type : Average

Compound	5.000	10.000	25.000	50.000	100.000	200.000	—	% RSD
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	RRF	—
99 n-Butanol	0.00708	0.00755	0.00826	0.00839	0.00871	0.00930	0.00822	9.712 <-
100 Methyl Methacrylate	0.17898	0.17553	0.18573	0.19952	0.20913	0.22296	0.19531	9.504
101 2-Nitropropane	0.05322	0.06193	0.06220	0.06218	0.06202	0.06322	0.06079	6.151
102 Chloropicrin	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
103 Cyclohexanone	0.02167	0.02329	0.02670	0.02927	0.03085	0.03125	0.02717	14.734
104 Pentachloroethane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
105 Benzyl Chloride	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
134 Thiophene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
135 Crotononitrile(1st Isomer)	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
136 Crotononitrile(2nd Isomer)	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
M 137 Total Crotononitrile	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
138 Paraldehyde	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
139 3,3,5-Trimethylcyclohexanone	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
140 1-Chlorohexane	+++++	+++++	+++++	+++++	+++++	+++++	0.76231	7.986
141 1,3,5-Trichlorobenzene	0.85746	0.79676	0.67661	0.75898	0.73463	0.74943	0.76231	5.658
143 Methyl Acetate	0.22313	0.21756	0.20107	0.19563	0.19918	0.19750	0.20568	13.218
144 Methylcyclohexane	0.34643	0.31604	0.23920	0.34078	0.28293	0.32667	0.30868	
145 Dimethoxymethane	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
146 2-Methylnaphthalene	+++++	+++++	+++++	+++++	+++++	+++++	+++++	+++++ <-
\$ 4 Dibromofluoromethane	0.22899	0.23170	0.22899	0.23960	0.23428	0.23320	0.23279	1.705
\$ 5 1,2-Dichloroethane-d4	0.31268	0.30905	0.31321	0.31488	0.33897	0.33187	0.32011	3.817
\$ 6 Toluene-d8	1.15061	1.17656	1.24969	1.21381	1.20619	1.21032	1.20120	2.832
\$ 7 Bromofluorobenzene	0.48378	0.51078	0.51438	0.50777	0.52150	0.52405	0.51038	2.827

STL North Canton

INITIAL CALIBRATION DATA

```

art Cal Date   : 16-AUG-2004 16:18
ld Cal Date   : 14-SEP-2004 15:41
lant Method   : ISTD
rgt Version   : 4.04
ntegrator     : HP RTE
ethod file    : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\82601LIUX11.m
l Date        : 14-Sep-2004 16:57 tapsvc

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libration File Names:
evel 1: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23875.D
evel 2: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23873.D
evel 3: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23872.D
evel 4: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23871.D
evel 5: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23870.D
evel 6: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23870.D

```

Compound	5.0000	10.0000	25.0000	50.0000	100.0000	200.0000	Curve	Coefficients	%RSD or R^2
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	b		
8 Dichlorodifluoromethane	0.28836	0.23754	0.28173	0.25561	0.24515	0.26401	AVRG	0.26140	7.79645
9 Chloromethane	0.55674	0.49898	0.45483	0.43975	0.42154	0.42948	AVRG	0.46672	11.13244
10 Vinyl Chloride	0.32349	0.31847	0.33246	0.31810	0.29804	0.3125	AVRG	0.31925	4.05579
11 Bromomethane	0.17259	0.16446	0.14190	0.15123	0.13819	0.14020	AVRG	0.15143	9.38575
12 Chloroethane	0.23244	0.24693	0.24362	0.23419	0.22089	0.22669	AVRG	0.23463	4.04335
13 Trichlorofluoromethane	0.38407	0.31147	0.34717	0.32533	0.30602	0.33365	AVRG	0.33462	8.49945
14 Dichlorofluoromethane	0.45871	0.50000	0.48561	0.49154	0.48286	0.49097	AVRG	0.48495	2.91496
15 Acrolein	0.03199	0.03065	0.03129	0.03093	0.03156	0.03098	AVRG	0.03123	1.55147
16 Acetone	72580	128042	262942	530389	973885	1907733	QUAD	-0.10527	0.81620

STL North Canton

INITIAL CALIBRATION DATA

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art Cal Date : 16-AUG-2004 16:18
d Cal Date : 14-SEP-2004 15:41
lant Method : ISTD
rget Version : 4.04
ntegrator : HP RTE
ethod file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\82601LUX11.m
,1 Date : 14-Sep-2004 16:57 tapsvc

```

Compound	5.000	10.000	25.000	50.000	100.000	200.000	curve	b	Coefficients	%RSD or R^2
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6			m1	
17 1,1-Dichloroethene	0.24613	0.23009	0.19938	0.23265	0.20714	0.22623 AVRG	-0. 02165	7.57113	-1.97871	0.99415
18 Freon-113	37991	76261	126246	390592	619208	1494012 QUAD	0.333050	3.62742		
19 Iodomethane	0.31149	0.34851	0.32631	0.33354	0.33020	0.33284 AVRG	0.84395 AVRG	0.83525		7.35551
20 Carbon Disulfide	0.91244	0.87319	0.74270	0.85398	0.78522	0.84395 AVRG	-0.19662	3.80728	-0.05949	0.99993
21 Methylene Chloride	169409	251128	413458	741159	1333682	2622600 QUAD	0.02976	0.02967		7.78866
22 Acetonitrile	0.03290	0.03019	0.03062	0.02865	0.02976	0.02552 AVRG	0.09022 AVRG	0.09228		2.15289
23 Acrylonitrile	0.9137	0.03573	0.09318	0.09088	0.09229	0.09022 AVRG	0.73394 AVRG	0.71686		5.70257
24 Methyl teri-butyl ether	0.65599	0.67280	0.74332	0.74432	0.74979	0.74979 AVRG	0.25710 AVRG	0.26526		4.43506
25 trans-1,2-Dichloroethene	0.28262	0.27489	0.25328	0.26749	0.25617	0.25617 AVRG	-0.00997	24.38425	-20.19946	0.99631
26 Hexane	11663	22298	38256	115096	195007	460062 QUAD	0.45290 AVRG	0.42446		6.03292
27 Vinyl acetate	0.41519	0.38277	0.42613	0.42027	0.44948	0.45290 AVRG	0.48389	3.75339		
28 1,1-Dichloroethane	0.51364	0.49348	0.46074	0.48352	0.47362	0.47833 AVRG	0.01945	5.26222		
29 tert-Butyl Alcohol	0.02052	0.02018	0.01934	0.01924	0.01980	0.01761 AVRG	0.13695	8.28054		
30 2-Butanone	0.15894	0.12780	0.13677	0.13776	0.13540	0.12902 AVRG	0.27198	3.88557		
M 31 1,2-Dichloroethene (total)	0.28520	0.28491	0.26194	0.27101	0.26404	0.26479 AVRG	0.27870	3.63568		
32 cis-1,2-dichloroethene	0.28778	0.29492	0.27453	0.27191	0.27248 AVRG	0.29523 AVRG	0.29161	3.90723		
33 2,2-Dichloropropane	0.29593	0.30774	0.27673	0.29383	0.28018					

Start Date : 14-Sep-2004 17:05

STL North Canton

INITIAL CALIBRATION DATA

```

Start Cal Date : 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Int Method : ISTD
Int Version : 4.04
egrator File : HP RTE
Method File : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-1C.b\8260LLUX11.m
1 Date : 14-Sep-2004 16:57 tapsvc

```

Compound	5.0000	10.0000	25.0000	50.0000	100.0000	200.0000	Curve	b	Coefficients	m1	m2	%RSD or R^2
	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6						
34 Bromochloromethane	0.12769	0.14200	0.12631	0.12865	0.12580	0.12308 AVRG		0.12905	0.48643	3.63104	5.07449	
35 Chloroform	0.51012	0.50575	0.47005	0.48650	0.47366	0.47250 AVRG		0.466039	0.99940	0.99940	0.99940	
36 Tetrahydrofuran	0.22560	0.32881	0.32881	0.32881	0.32881	0.32912 AVRG		12.83982	4.66039	5.87801	5.87801	
37 1,1,1-Trichloroethane	0.39287	0.37451	0.30656	0.35792	0.32912	0.30261 AVRG		0.36737	0.36737	6.00098	6.00098	
38 1,1-Dichloropropene	0.30286	0.29259	0.24537	0.31080	0.27649	0.31080 AVRG		0.39169	0.39169	5.46736	5.46736	
39 Carbon Tetrachloride	0.40025	0.42402	0.38489	0.39237	0.39237	0.39237 AVRG		1.10254 AVRG	1.10254 AVRG	1.15116	2.81819	
40 1,1,2-Dichloroethane	1.25995	1.19011	1.13415	1.13415	1.12471	1.12471 AVRG		0.26258 AVRG	0.26258 AVRG	0.26272	4.24139	
41 Benzene	0.27283	0.26390	0.25129	0.26732	0.25129	0.26732 AVRG		0.27607 AVRG	0.27607 AVRG	0.28397	8.90709 <-	
42 Trichloroethylene	0.29669	0.30122	0.27194	0.28098	0.27691	0.28098 AVRG		0.00260	0.00260	0.00243	3.33962	
43 1,2-Dichloropropane	0.00242	0.00233	0.00258	0.00261	0.00261	0.00261 AVRG		0.15545 AVRG	0.15545 AVRG	0.15736	0.15736	
44 1,4-Dioxane	0.15285	0.16666	0.15233	0.15822	0.15822	0.15822 AVRG		0.37355	0.37355	2.34541	2.34541	
45 Dibromomethane	0.38193	0.38487	0.36100	0.37042	0.36100	0.37042 AVRG		0.36931	0.36931	0.37381 AVRG	8.38480	
46 Bromodichloromethane	0.13950	0.14798	0.15985	0.16693	0.17284	0.17092 AVRG		0.17092 AVRG	0.17092 AVRG	0.15967	4.05753	
47 2-Chloroethyl vinyl ether	0.43417	0.47734	0.43633	0.45487	0.46634	0.47331 AVRG		0.47331 AVRG	0.47331 AVRG	0.45706	3.42375	
48 cis-1,3-Dichloropropene	0.23661	0.23635	0.24230	0.25213	0.25538	0.25182 AVRG		0.25538	0.25538	1.49327	1.98694	
49 4-Methyl-2-pentanone	1.48717	1.54897	1.47049	1.49192	1.47049	1.47049 AVRG		1.47049 AVRG	1.47049 AVRG	1.49327	1.98694	
50 Toluene												

STL North Canton

INITIAL CALIBRATION DATA

```

:art Cal Date   : 16-AUG-2004 16:18
:id Cal Date   : 14-SEP-2004 15:41
:rant Method   : ISTD
:rgent Version  : 4.04
:tegrator      : HP RTE
:thod file     : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
:ll Date       : 14-Sep-2004 16:57 tapsvc

```

Compound	5.0000	10.0000	25.0000	50.0000	100.0000	200.0000	curve	b	Coefficients	m1	m2	%RSD or R^2
	Level 1	Level 2	Level 3	level 4	Level 5	Level 6						
51 trans-1,3-Dichloropropene	0.50647	0.53118	0.51429	0.54236	0.56606	0.56198 AVRG		0.53539		4.16176		
52 Ethyl Methacrylate	0.38934	0.45025	0.45000	0.4861	0.49453	0.49921 AVRG		0.46066		8.88545		
53 1,1,2-Trichloroethane	0.30188	0.31382	0.29748	0.30352	0.30279	0.29789 AVRG		0.30290		1.95346		
54 1,3-Dichloropropane	0.57206	0.59182	0.56392	0.55915	0.57182	0.56489 AVRG		0.57071		1.99579		
55 Tetrachloroethene	0.25509	0.26206	0.22142	0.2459	0.22718	0.24073 AVRG		0.24193		6.47809		
56 2-Hexanone	0.22234	0.23807	0.24231	0.24889	0.24506	0.25308 AVRG		0.24179		4.48622		
57 Dibromochloromethane	0.30967	0.33372	0.31997	0.31951	0.32624	0.32930 AVRG		0.32307		2.64150		
58 1,2-Dibromoethane	0.26921	0.31637	0.29041	0.30230	0.30732	0.30501 AVRG		0.29844		5.56079		
59 Chlorobenzene	0.96525	1.03980	0.94396	0.94705	0.94928	0.95592 AVRG		0.9688		3.77675		
60 1,1,1,2-Tetrachloroethane	0.33779	0.34414	0.32638	0.32925	0.34031	0.33757 AVRG		0.33590		2.01341		
61 Ethylbenzene	0.46024	0.50768	0.46972	0.48906	0.48935	0.51358 AVRG		0.48827		4.24550		
62 m + p-Xylene	0.60709	0.64675	0.61837	0.63926	0.62027	0.64155 AVRG		0.62888		2.51086		
63 Xylenes (total)	0.59346	0.64358	0.61984	0.63276	0.62064	0.64001 AVRG		0.62505		2.92467		
64 Xylene-o	0.56221	0.63725	0.62277	0.61975	0.62137	0.63694 AVRG		0.61738		4.25246		
65 Styrene	1.00309	1.11779	1.08299	1.11462	1.13838	1.16736 AVRG		1.10404		5.14327		
66 Bromoform	0.20132	0.22491	0.21511	0.21840	0.22531	0.23254 AVRG		0.21960		4.91986		
67 Isopropylbenzene	1.34498	1.36235	1.30493	1.41191	1.35691	1.46909 AVRG		1.37503		4.17959		

STL North Canton

INITIAL CALIBRATION DATA

```

art Cal Date : 16-AUG-2004 16:18
d Cal Date : 14-SEP-2004 15:41
ant Method : ISTD
rget Version : 4.04
tegrator : HP RTE
thod file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-1C.b\8260LLUX11.m
1 Date : 14-Sep-2004 16:57 tapsvc

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Compound	Level 1	Level 2	Level 3	Level 4	Level 5	Curve	b	Coefficients	%RSD
							ml	m2	or R^2
	5.0000	10.0000	25.0000	50.0000	100.0000	200.0000			
68 1,1,2,2-Tetrachloroethane	0.74074	0.78253	0.73952	0.75703	0.76116	0.74128 AVRG		0.75371	2.23655
69 1,4-Dichloro-2-butene	0.23161	0.23946	0.23911	0.25172	0.26183	0.26419 AVRG		0.24799	5.3736
70 1,2,3-Trichloropropane	0.25205	0.24157	0.24830	0.24773	0.24135	0.24135 AVRG		0.24733	2.01646
71 Bromobenzene	0.74497	0.77908	0.72155	0.75186	0.74513	0.73336 AVRG		0.74599	2.60484
72 n-Propylbenzene	0.71725	0.70753	0.62905	0.72290	0.67589	0.72046 AVRG		0.68879	2.88741
73 2-Chlorotoluene	0.71622	0.69234	0.65887	0.70074	0.67558	0.68895 AVRG		2.31439	3.87363
74 1,3,5-Trimethylbenzene	2.25968	2.26774	2.20534	2.39411	2.31559	2.44388 AVRG		0.73580	3.30376
75 4-Chlorotoluene	0.75654	0.75820	0.70598	0.74482	0.71695	0.72231 AVRG		0.87499	5.07853
76 tert-Butylbenzene	1.89049	1.94223	1.70074	1.94123	1.83516	1.94010 AVRG		4.49215	
77 1,2,4-Trimethylbenzene	2.33163	2.49724	2.38803	2.57507	2.50096	2.58086 AVRG		2.47063	5.80556
78 sec-Butylbenzene	2.66025	2.55624	2.29546	2.65769	2.44749	2.62040 AVRG		2.53792	
79 4-Isopropyltoluene	2.02816	2.10772	1.90515	2.27140	2.10588	2.25071 AVRG		2.11151	6.51441
80 1,3-Dichlorobenzene	1.50394	1.41628	1.30596	1.36771	1.33637	1.33353 AVRG		1.37723	5.27600
81 1,4-Dichlorobenzene	1.49216	1.50657	1.37893	1.45011	1.40876	1.40879 AVRG		1.44089	3.5234
82 n-Butylbenzene	1.86201	1.92154	1.64744	2.01989	1.84399	2.02271 AVRG		1.88626	7.3890
83 1,2-Dichlorobenzene	1.44948	1.40634	1.30068	1.35160	1.32364	1.30712 AVRG		1.35648	4.3970
84 1,2-Dibromo-3-chloropropane	0.12641	0.12593	0.12612	0.12831	0.13070	0.12832 AVRG		0.12763	1.4461

STL North Canton

INITIAL CALIBRATION DATA

```

:part Cal Date : 16-AUG-2004 16:18
:id Cal Date : 14-SEP-2004 15:41
:rant Method : ISTD
:arget Version : 4.04
:ntegrator : HP RTE
:stethod file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\82601LUX11.m
:al Date : 14-Sep-2004 16:57 tapsvc

```

Compound	5.0000 Level 1	10.0000 Level 2	25.0000 Level 3	50.0000 Level 4	100.0000 Level 5	200.0000 Level 6	Curve 	b 	Coefficients m1 	%RSD or R^2
85 1,2,4-Trichlorobenzene	0.56391	0.60976	0.52246	0.58135	0.57721	0.60830 AVRG		0.57750	5.59592	
86 Hexachlorobutadiene	36648	62649	126348	272701	475339	979903 QUAD	-0.05586	4.17449	0.99883	
87 Naphthalene	1.19572	1.26888	1.21865	1.39469	1.53442	1.56161 AVRG		1.33300	11.77993	
88 1,2,3-Trichlorobenzene	0.38318	0.44192	0.36462	0.41931	0.44173	0.43401 AVRG		0.41413	7.90998	
89 Ethyl Ether	0.26862	0.25572	0.24190	0.23651	0.23958	0.23653 AVRG		0.24654	5.24098	
90 Ethanol	+***+	+***+	+***+	+***+	+***+	+***+ AVRG		0.000e+000	0.000e+000 <-	
91 3-Chloropropene	0.08501	0.09752	0.10420	0.10696	0.10892	0.11570 AVRG		0.10305	10.33497	
92 Isopropyl Ether	0.19894	0.21144	0.22424	0.23464	0.23633	0.23561 AVRG		0.22353	6.88556	
93 2-Chloro-1,3-butadiene	0.33010	0.35251	0.38261	0.39081	0.38671	0.39380 AVRG		0.37276	6.88550	
94 Propionitrile	0.04383	0.04396	0.04410	0.04111	0.04094	0.03930 AVRG		0.04231	4.40359	
95 Ethyl Acetate	0.25153	0.23918	0.24678	0.23828	0.24450	0.25022 AVRG		0.24508	2.25277	
96 Methacrylonitrile	0.15470	0.15983	0.15530	0.15862	0.16298	0.16195 AVRG		0.15890	2.13440	
97 Isobutanol	0.01214	0.01067	0.01151	0.01109	0.01159	0.01152 AVRG		0.01142	4.36240	
98 Cyclohexane	0.37486	0.38746	0.29089	0.40866	0.34684	0.39833 AVRG		0.36784	11.78329	
99 n-Butanol	0.00708	0.00755	0.00826	0.00839	0.00871	0.00930 AVRG		0.00822	9.71167 <-	
100 Methyl Methacrylate	0.17898	0.17553	0.18573	0.19952	0.20913	0.22296 AVRG		0.19531	9.50417	
101 2-Nitropropane	0.05322	0.06193	0.06220	0.06218	0.06202	0.06322 AVRG		0.06079	6.15064	

Date : 14-Sep-2004 17:05

STL North Canton

INITIAL CALIBRATION DATA

1st Cal Date : 16-AUG-2004 16:18
 1st Cal Date : 14-SEP-2004 15:41
 Int Method : ISTD
 Target Version : 4.04
 Integrator : HP RTE
 Method file : \\gcanoh04\dd\chem\MSV\aux11.i\J40914B-IC.b\8260LIUX11.m
 1st Date : 14-Sep-2004 16:57 tapsvc

Compound	Coefficients						%RSD or R^2						
	m1	m2	b	Curve	Level 1	Level 2							
				5.0000	10.0000	25.0000	50.0000	100.0000	200.0000	Level 4	Level 5	Level 6	
102 Chloropicrin	0.02167	0.02329	0.02670	+++++	0.02927	0.03085	0.03125	AVRG	+++++	0.000e+000	0.02717	0.000e+000	
103 Cyclohexanone				+++++									
104 Pentachloroethane				+++++									
105 Benzyl Chloride				+++++									
134 Thiophene				+++++									
135 Crotononitrile(1st Isomer)				+++++									
136 Crotononitrile(2nd Isomer)				+++++									
1 137 Total Crotononitrile				+++++									
138 Paraldehyde				+++++									
139 3,3,5-Trimethylcyclohexanone				+++++									
140 1-Chlorohexane	0.85746	0.79676	0.67661	+++++	0.7598	0.73463	0.74943	AVRG	+++++	0.000e+000	0.20568	0.000e+000	
141 1,3,5-Trichlorobenzene	0.22313	0.21756	0.20107	+++++	0.19663	0.19318	0.19750	AVRG	+++++	0.000e+000	0.30868	0.000e+000	
143 Methyl Acetate	0.34643	0.31604	0.23920	+++++	0.34078	0.28293	0.32667	AVRG	+++++	0.000e+000	0.000e+000	0.000e+000	
144 Methylcyclohexane				+++++									
145 Dimethylnaphthalene				+++++									
146 2-Methylnaphthalene				+++++									

port Date : 14-Sep-2004 17:05

STL North Canton

INITIAL CALIBRATION DATA

```

art Cal Date : 16-AUG-2004 16:18
id Cal Date : 14-SEP-2004 15:41
lant Method : ISTD
rgt Version : 4.04
tegrator : HP RTE
ethod file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260ILLUX11.m
il Date : 14-Sep-2004 16:57 tapsvc

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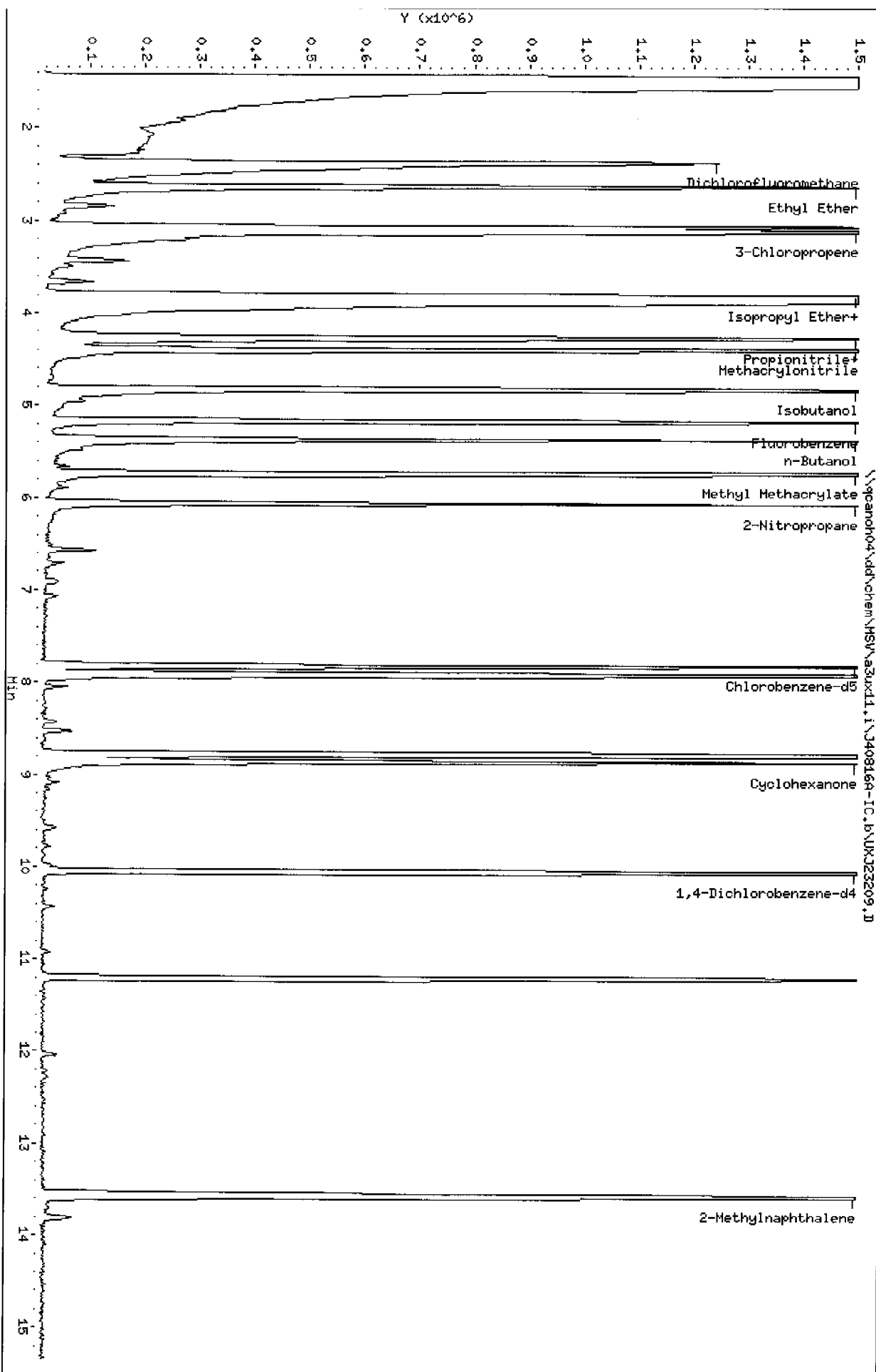
Compound	Level 1	Level 2	Level 3	Level 4	Level 5	Level 6	Curve	b	Coefficients	%RSD
							m1	m2	or R^2	
4 Dibromofluoromethane	0.22899	0.23170	0.22899	0.23960	0.23428	0.23320	AVRG	0.23279	1.70494	
5 1,2-Dichloroethane-d4	0.31268	0.30905	0.31321	0.31481	0.33897	0.33187	AVRG	0.32011	3.81724	
6 Toluene-d8	1.15061	1.17656	1.24959	1.241381	1.20619	1.21032	AVRG	1.20120	2.83165	
7 Bromofluorobenzene	0.48378	0.51078	0.51438	0.50777	0.52150	0.52405	AVRG	0.51038	2.83658	

Curve	Formula	Units
Averaged	Amt = Rsp*m1	Response
Quad	Amt = b + m1*Rsp + m2*Rsp^2	Response

Data File: \\qcanoh04\dd\chem\MSV\aa3ux11.i\J40816A-IC.b\UXJ23209.II

Sample Info: 200NG-ASIC
Purge Volume: 5.0
Column phase: DB624

Operator: 43582
Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23209.D
Report Date: 17-Aug-2004 14:56

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23209.D
Lab Smp Id: 200NG-A9IC
Inj Date : 16-AUG-2004 16:18
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 200NG-A9IC
Misc Info : J40816A-IC,8260LLUX11,3-IX.SUB,43582,1,6
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:56 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 8 Calibration Sample, Level: 6
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	2040512	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1444382	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	767740	50.0000		
14 Dichlorofluoromethane	67	2.367	2.367 (0.458)	4007314	200.000	202.48 (A)	
89 Ethyl Ether	59	2.627	2.627 (0.508)	1930594	200.000	191.88	
91 3-Chloropropene	76	3.112	3.112 (0.602)	944322	200.000	224.54 (A)	
92 Isopropyl Ether	87	3.810	3.810 (0.737)	9615196	1000.00	1054.0 (A)	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	3214209	200.000	211.29 (A)	
94 Propionitrile	54	4.260	4.260 (0.824)	651311	400.000	377.22 (A)	
95 Ethyl Acetate	43	4.260	4.260 (0.824)	4084685	400.000	408.39 (A)	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	1321846	200.000	203.84 (A)	
97 Isobutanol	41	4.816	4.816 (0.616)	1331366	4000.00	4036.3 (A)	
99 n-Butanol	56	5.361	5.361 (0.685)	1074873	4000.00	4527.6 (A)	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	1819805	200.000	228.32 (A)	
101 2-Nitropropane	41	6.059	6.059 (1.172)	1031981	400.000	415.94 (A)	
103 Cyclohexanone	55	8.851	8.851 (0.881)	959642	2000.00	2300.0 (A)	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	4079814	400.000	1085.6 (A)	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23209.D
Report Date: 17-Aug-2004 14:56

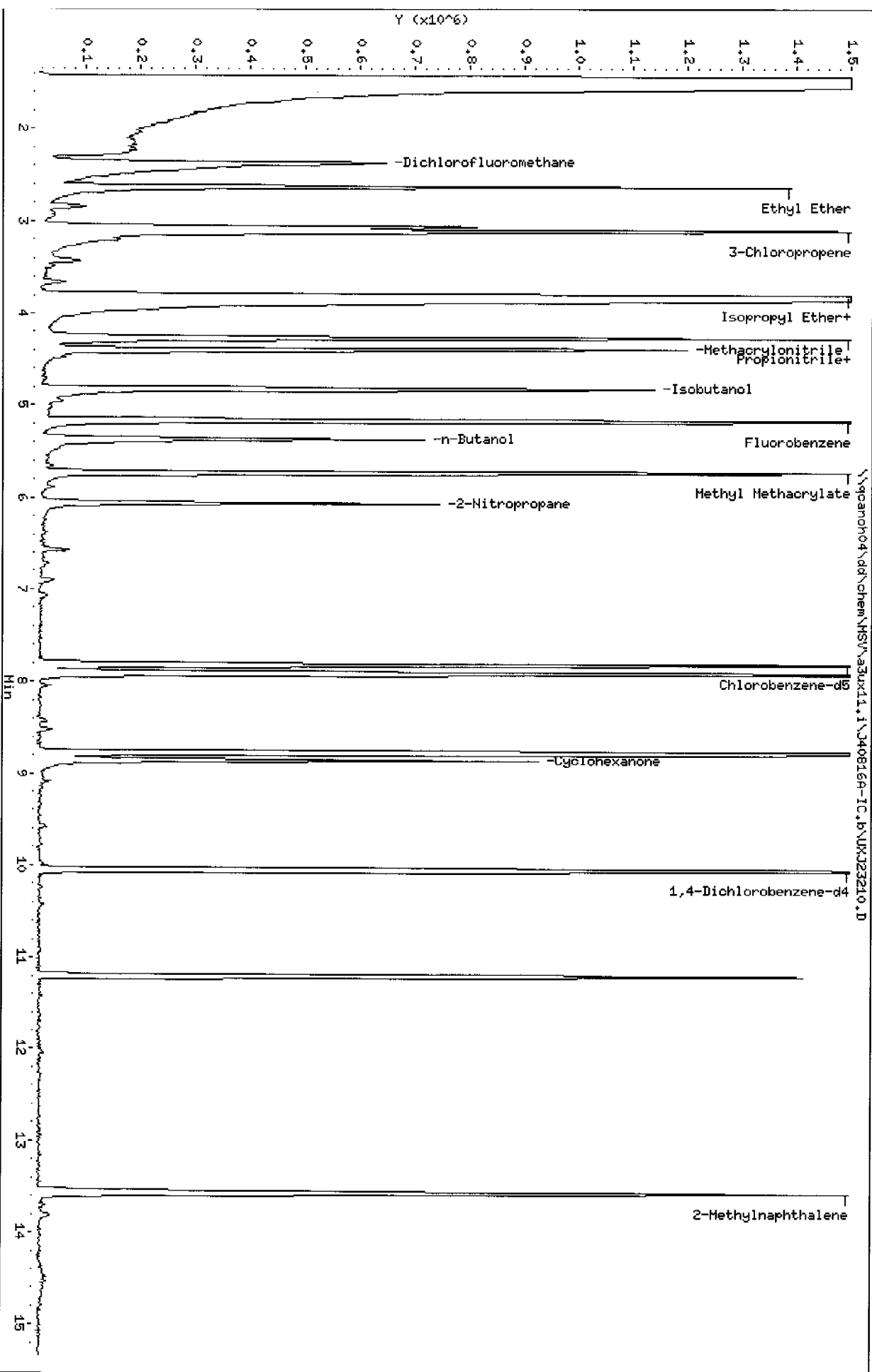
QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Purge Volume: 5.0
Column phase: DB624

Instrument: a30x11.i

Operator: 43582
Column diameter: 0.18



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\UXJ23210.D
Report Date: 17-Aug-2004 14:57

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\UXJ23210.D
Lab Smp Id: 100NG-A9IC
Inj Date : 16-AUG-2004 16:40
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 100NG-A9IC
Misc Info : J40816A-IC,8260LLUX11,3-IX.SUB,43582,1,5
Comment :
Method : \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\8260LLUX11.m
Meth Date : 17-Aug-2004 14:56 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 9 Calibration Sample, Level: 5
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	1987706	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1429041	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	734335	50.0000		
14 Dichlorofluoromethane	67	2.367	2.367 (0.458)	1919583	100.000	99.570	
89 Ethyl Ether	59	2.627	2.627 (0.508)	952411	100.000	97.174	
91 3-Chloropropene	76	3.112	3.112 (0.602)	432982	100.000	105.69	
92 Isopropyl Ether	87	3.810	3.810 (0.737)	4697492	500.000	528.62 (A)	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	1537316	100.000	103.74	
94 Propionitrile	54	4.260	4.260 (0.824)	325535	200.000	193.55	
95 Ethyl Acetate	43	4.260	4.260 (0.824)	1943985	200.000	199.52	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	647907	100.000	102.57	
97 Isobutanol	41	4.816	4.816 (0.616)	662388	2000.00	2029.7 (A)	
99 n-Butanol	56	5.361	5.361 (0.685)	498130	2000.00	2120.8 (A)	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	831362	100.000	107.08	
101 2-Nitropropane	41	6.059	6.059 (1.172)	493098	200.000	204.02 (A)	
103 Cyclohexanone	55	8.851	8.851 (0.881)	453132	1000.00	1135.4 (A)	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	1299383	200.000	361.47 (A)	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23210.D
Report Date: 17-Aug-2004 14:57

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\pcanoh04\\dd\\chem\\HSW\\a30x11.i\\J40816A-IC.b\\UXJ23211.D
Date : 16-AUG-2004 17:03
Client ID:
Sample Info: 50HG-APIC

Purge Volume: 5.0
Column phase: DB624

Instrument: a30x11.i

Operator: 43582

Column diameter: 0.18

1.5

1.4

1.3

1.2

1.1

1.0

0.9

0.8

0.7

0.6

0.5

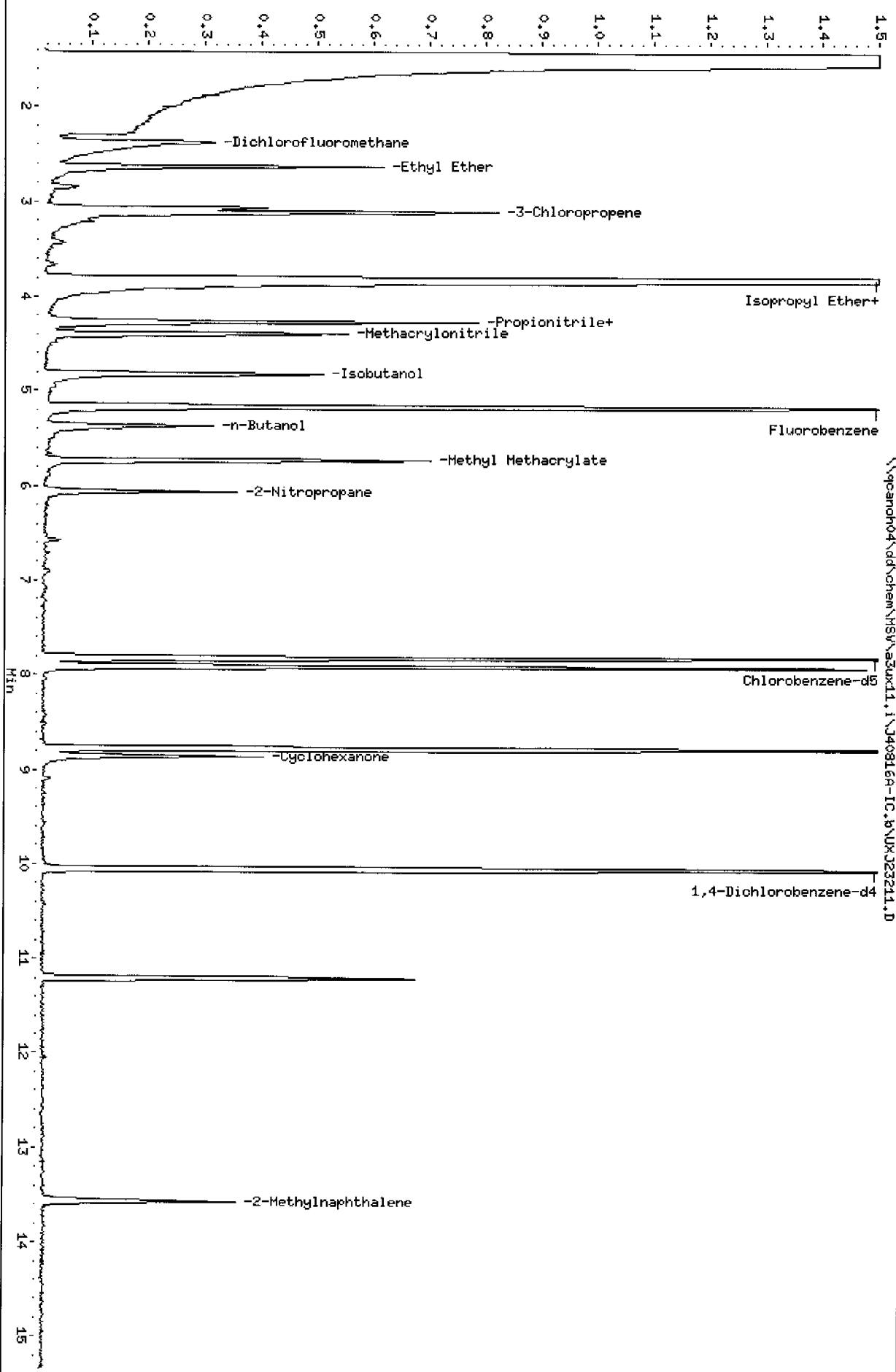
0.4

0.3

0.2

0.1

Y ($\times 10^6$)



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23211.D
Lab Smp Id: 50NG-A9IC
Inj Date : 16-AUG-2004 17:03
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 50NG-A9IC
Misc Info : J40816A-IC,8260LLUX11,3-IX.SUB,43582,1,4
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:57 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 10 Calibration Sample, Level: 4
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	1946935	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1394264	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	698314	50.0000		
14 Dichlorofluoromethane	67	2.379	2.379 (0.460)	957001	50.0000	50.680	
89 Ethyl Ether	59	2.639	2.639 (0.510)	461257	50.0000	48.047	
91 3-Chloropropene	76	3.112	3.112 (0.602)	208248	50.0000	51.898	
92 Isopropyl Ether	87	3.810	3.810 (0.737)	2284128	250.000	262.42 (A)	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	760882	50.0000	52.422	
94 Propionitrile	54	4.260	4.260 (0.824)	160094	100.000	97.178	
95 Ethyl Acetate	43	4.272	4.272 (0.826)	927819	100.000	97.223	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	308829	50.0000	49.914	
97 Isobutanol	41	4.816	4.816 (0.616)	309158	1000.00	970.97 (A)	
99 n-Butanol	56	5.361	5.361 (0.685)	234037	1000.00	1021.2 (A)	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	388455	50.0000	51.079	
101 2-Nitropropane	41	6.059	6.059 (1.172)	242106	100.000	102.27	
103 Cyclohexanone	55	8.851	8.851 (0.881)	204421	500.000	538.66 (A)	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	247408	100.000	72.376	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23211.D
Report Date: 17-Aug-2004 14:57

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

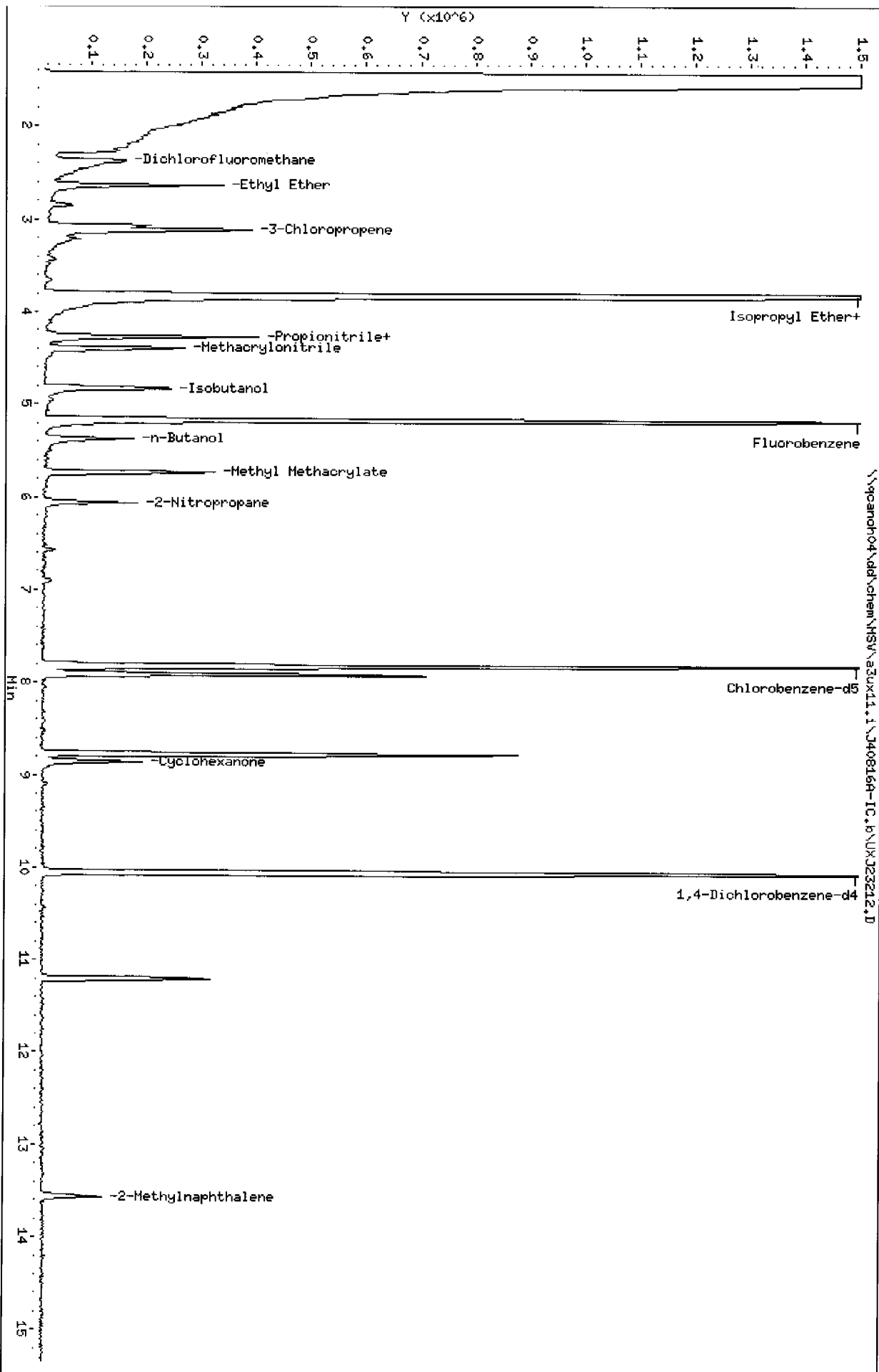
Client ID:
Sample Info: 25HG-A9IC
Purge Volume: 5.0

Column phase: DB624

Instrument: z30x11.i

Operator: 43582
Column diameter: 0.18

Y ($\times 10^6$)



Data File: \\qcanoh04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\UXJ23212.D
Report Date: 17-Aug-2004 14:58

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\UXJ23212.D
Lab Smp Id: 25NG-A9IC
Inj Date : 16-AUG-2004 17:26
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 25NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 3
Comment :
Method : \\QCANOH04\dd\chem\MSV\ a3ux11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:57 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 11 Calibration Sample, Level: 3
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	1894679	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1362249	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	685784	50.0000		
14 Dichlorofluoromethane	67	2.378	2.378 (0.460)	460035	25.0000	25.034	
89 Ethyl Ether	59	2.639	2.639 (0.510)	229157	25.0000	24.529	
91 3-Chloropropene	76	3.112	3.112 (0.602)	98714	25.0000	25.279	
92 Isopropyl Ether	87	3.810	3.810 (0.737)	1062158	125.000	125.40	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	362465	25.0000	25.661	
94 Propionitrile	54	4.260	4.260 (0.824)	83549	50.0000	52.114	
95 Ethyl Acetate	43	4.272	4.272 (0.826)	467574	50.0000	50.347	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	147119	25.0000	24.434	
97 Isobutanol	41	4.816	4.816 (0.616)	156732	500.000	503.82 (A)	
99 n-Butanol	56	5.372	5.372 (0.687)	112573	500.000	502.78 (A)	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	175947	25.0000	23.774	
101 2-Nitropropane	41	6.059	6.059 (1.172)	117850	50.0000	51.156	
103 Cyclohexanone	55	8.851	8.851 (0.881)	91550	250.000	245.65 (A)	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	80002	50.0000	23.831	

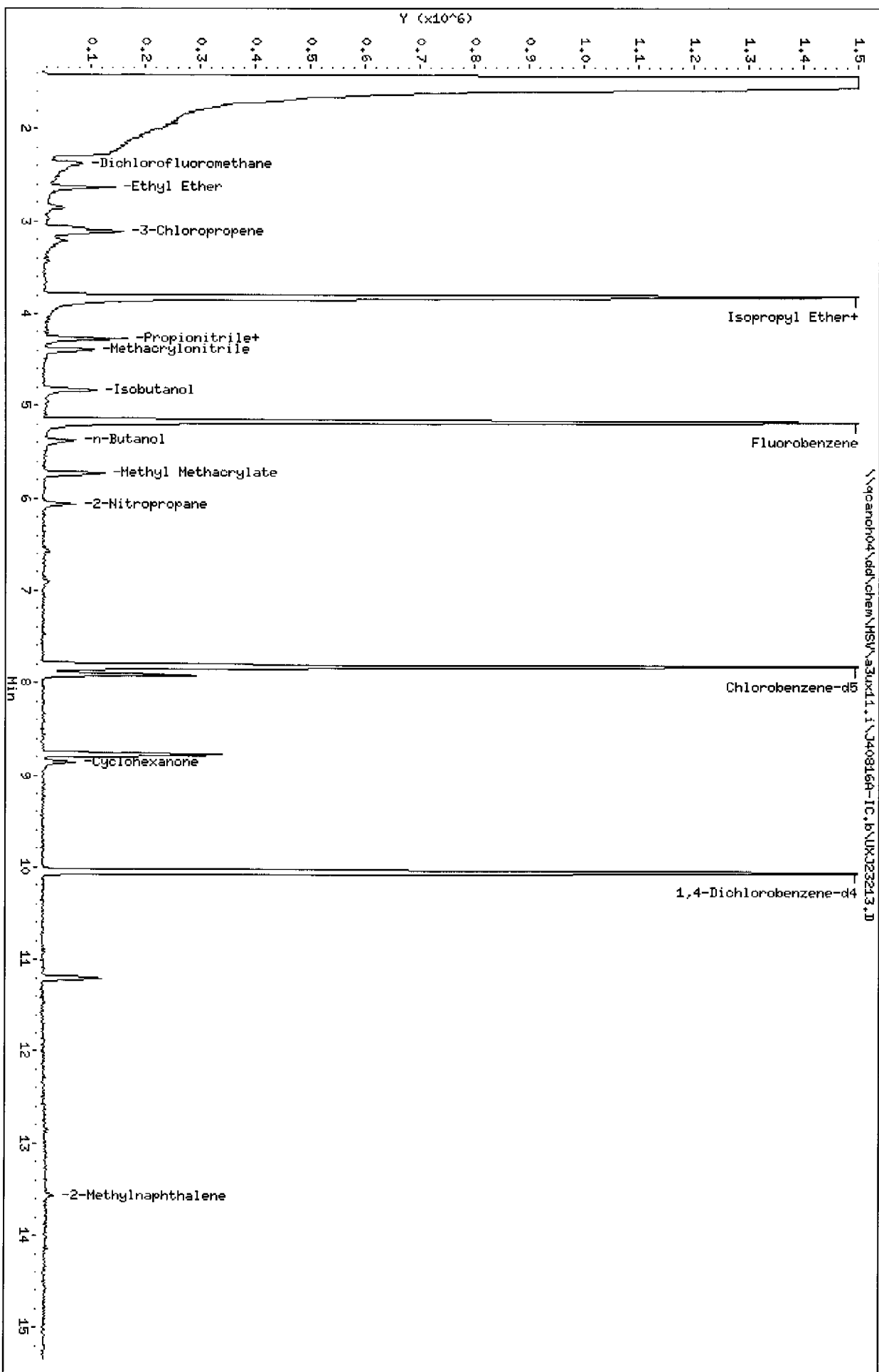
Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40816A-IC.b\UXJ23212.D
Report Date: 17-Aug-2004 14:58

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\qcancho04\\dd\\chem\\MSV\\a30x11.i\\J40816A-IC.b\\UKJ3213.D
Date : 16-AUG-2004 17:48
Client ID:
Sample Info: 10NG-A9IC
Purge Volume: 5.0
Column phase: DB624

Instrument: a30x11.i
Operator: 43582
Column diameter: 0.18
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Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\UXJ23213.D
Report Date: 17-Aug-2004 14:58

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\UXJ23213.D
Lab Smp Id: 10NG-A9IC
Inj Date : 16-AUG-2004 17:48
Operator : 43582 Inst ID: A3UX11.i
Smp Info : 10NG-A9IC
Misc Info : J40816A-IC, 8260LLUX11, 3-IX.SUB, 43582, 1, 2
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40816A-IC.b\8260LLUX11.m
Meth Date : 17-Aug-2004 14:58 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 12 Calibration Sample, Level: 2
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

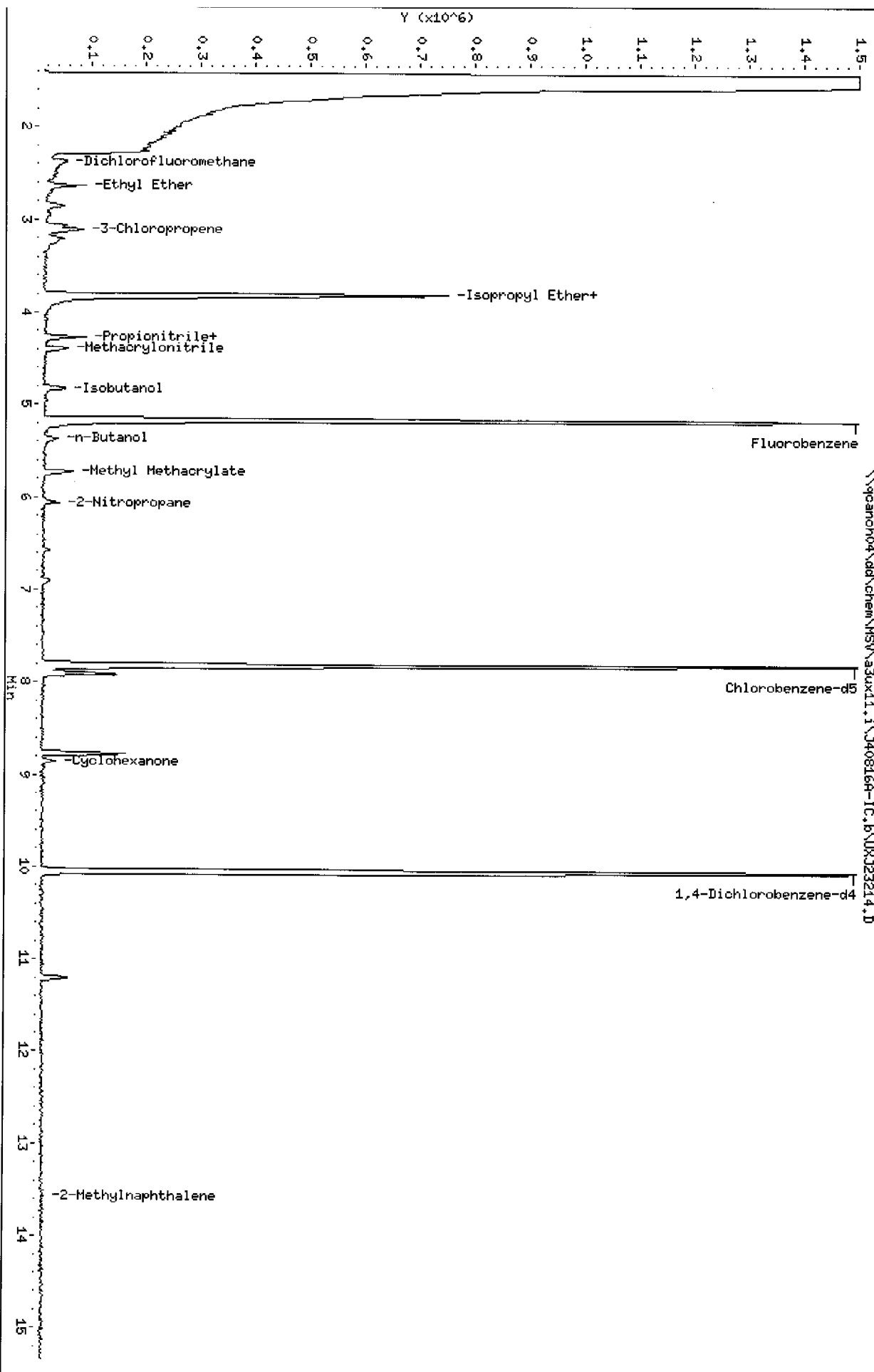
Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	1783996	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1312732	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	643754	50.0000		
14 Dichlorofluoromethane	67	2.379	2.379 (0.460)	178398	10.0000	10.310	
89 Ethyl Ether	59	2.639	2.639 (0.510)	91239	10.0000	10.372	
91 3-Chloropropene	76	3.112	3.112 (0.602)	34795	10.0000	9.463	
92 Isopropyl Ether	87	3.810	3.810 (0.737)	377202	50.0000	47.294	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	125775	10.0000	9.457	
94 Propionitrile	54	4.260	4.260 (0.824)	31372	20.0000	20.782	
95 Ethyl Acetate	43	4.272	4.272 (0.826)	170678	20.0000	19.518	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	57028	10.0000	10.059	
97 Isobutanol	41	4.828	4.828 (0.617)	56016	200.000	186.86	
99 n-Butanol	56	5.384	5.384 (0.688)	39656	200.000	183.79	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	62629	10.0000	8.987	
101 2-Nitropropane	41	6.059	6.059 (1.172)	44196	20.0000	20.375	
103 Cyclohexanone	55	8.851	8.851 (0.881)	29984	100.000	85.706	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	13721	20.0000	4.354	

Data File: \\qpcancho4\\dd\\Chem\\MSI\\a30x11.i\\J40816A-IC.b\\UXJ23214.D
Date : 16-AUG-2004 18:11
Client ID:
Sample Info: SNC-A91C
Purge Volume: 5.0
Column phase: DB624

Instrument: a30x11.i
Operator: 43582
Column diameter: 0.18



Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\UXJ23214.D
Report Date: 17-Aug-2004 14:59

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\UXJ23214.D
Lab Smp Id: 5NG-A9IC
Inj Date : 16-AUG-2004 18:11
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 5NG-A9IC
Misc Info : J40816A-IC,8260LLUX11,3-IX.SUB,43582,1,1
Comment :
Method : \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\8260LLUX11.m
Meth Date : 17-Aug-2004 14:59 evansl Quant Type: ISTD
Cal Date : 16-AUG-2004 18:11 Cal File: UXJ23214.D
Als bottle: 13 Calibration Sample, Level: 1
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

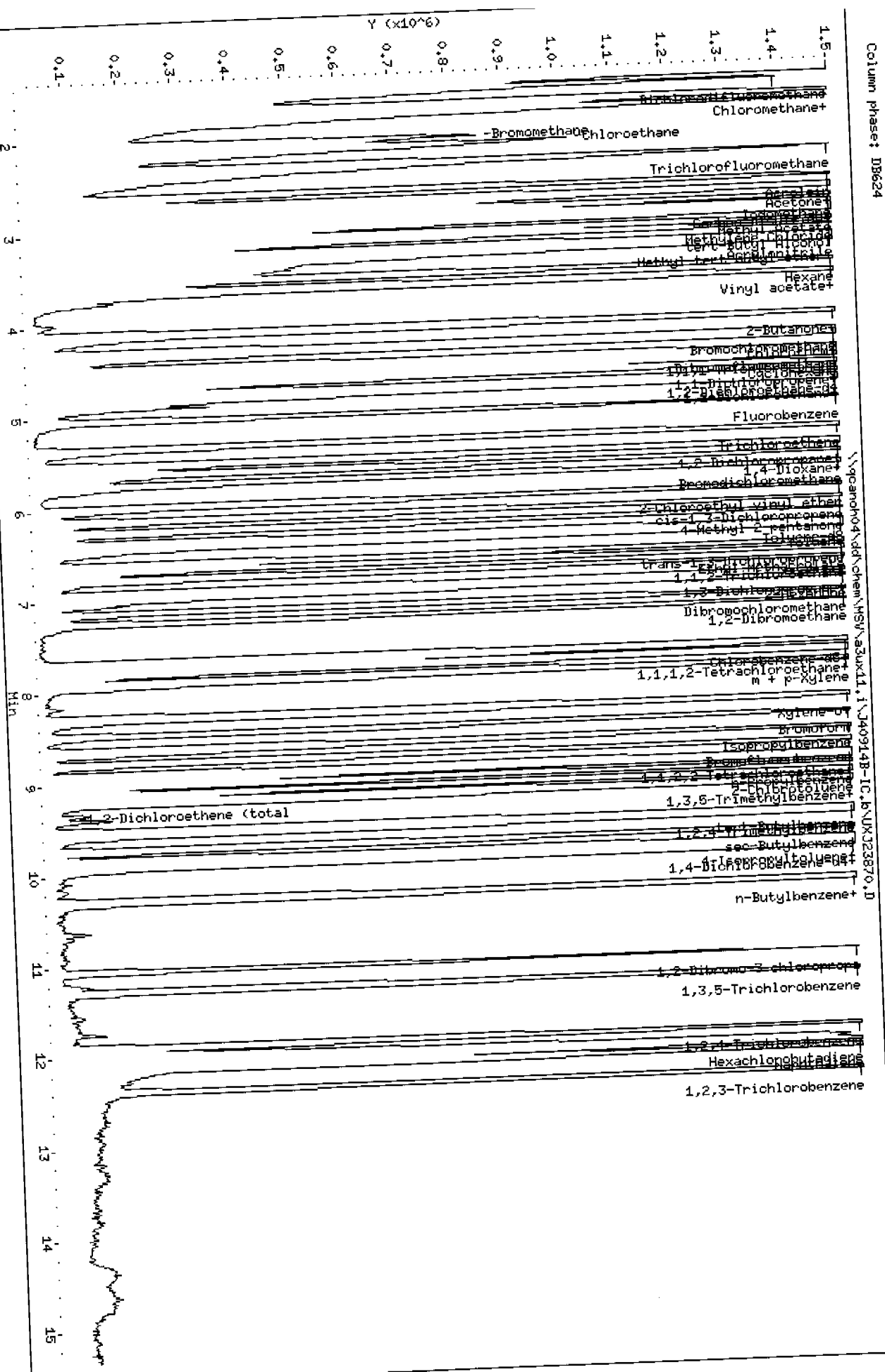
Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
* 1 Fluorobenzene	96	5.171	5.171 (1.000)	1729289	50.0000		
* 2 Chlorobenzene-d5	117	7.822	7.822 (1.000)	1295054	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	10.046	10.046 (1.000)	623017	50.0000		
14 Dichlorofluoromethane	67	2.379	2.379 (0.460)	79325	5.00000	4.730	
89 Ethyl Ether	59	2.639	2.639 (0.510)	46453	5.00000	5.448	
91 3-Chloropropene	76	3.112	3.112 (0.602)	14700	5.00000	4.124	
92 Isopropyl Ether	87	3.822	3.822 (0.739)	172012	25.0000	22.250	
93 2-Chloro-1,3-butadiene	53	3.846	3.846 (0.744)	57084	5.00000	4.428	
94 Propionitrile	54	4.260	4.260 (0.824)	15160	10.0000	10.360	
95 Ethyl Acetate	43	4.272	4.272 (0.826)	86994	10.0000	10.263	
96 Methacrylonitrile	41	4.390	4.390 (0.849)	26752	5.00000	4.868	
97 Isobutanol	41	4.816	4.816 (0.616)	31443	100.000	106.32	
99 n-Butanol	56	5.372	5.372 (0.687)	18347	100.000	86.193	
100 Methyl Methacrylate	41	5.727	5.727 (1.108)	30951	5.00000	4.582	
101 2-Nitropropane	41	6.059	6.059 (1.172)	18407	10.0000	8.754	
103 Cyclohexanone	55	8.851	8.851 (0.881)	13502	50.0000	39.878	
146 2-Methylnaphthalene	142	13.561	13.561 (1.350)	1849	10.0000	0.6063	

Data File: \\qcanoh04\\dd\\chem\\MSV\\a3Ux11.i\\J40914B-IC.b\\UX323870.D
 Date : 14-SEP-2004 13:48
 Client ID:
 Sample Info: 200NG-IC
 Purge Volume: 5.0
 Column Phase: IB624

Instrument: a3Ux11.i

Operator: 43582
 Column diameter: 0.18
 \\qcanoh04\\dd\\chem\\MSV\\a3Ux11.i\\J40914B-IC.b\\UX323870.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23870.D
Report Date: 15-Sep-2004 12:45

STL North Canton

VOLATILE REPORT SW-846 Method

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23870.D
Lab Smp Id: 200NG-IC
Inj Date : 14-SEP-2004 13:48 Inst ID: a3ux11.i
Operator : 43582
Smp Info : 200NG-IC
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,1,6
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m Quant Type: ISTD
Meth Date : 15-Sep-2004 12:45 evans1 Cal File: UXJ23875.D
Cal Date : 14-SEP-2004 15:41 Calibration Sample, Level: 6
Als bottle: 2
Dil Factor: 1.00000 Compound Sublist: 2-8260.SUB
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT	ON-COL
							(ng)	(ng)
*	1 Fluorobenzene	96	5.029	5.029 (1.000)		2337325	50.0000	
*	2 Chlorobenzene-d5	117	7.668	7.668 (1.000)		1845289	50.0000	
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)		1029809	50.0000	
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.889)		2180284	200.000	200.35(A)
\$	5 1,2-Dichloroethane-d4	65	4.745	4.745 (0.944)		3102729	200.000	207.35(A)
\$	6 Toluene-d8	98	6.366	6.366 (0.830)		8933577	200.000	201.52(A)
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.145)		3868072	200.000	205.36(A)
\$	8 Dichlorodifluoromethane	85	1.527	1.527 (0.304)		2468303	200.000	202.00(A)
9	Chloromethane	50	1.680	1.680 (0.334)		4015293	200.000	184.04
10	Vinyl Chloride	62	1.763	1.763 (0.351)		3096925	200.000	207.51(A)
11	Bromomethane	94	2.047	2.047 (0.407)		1310746	200.000	185.17
12	Chloroethane	64	2.118	2.118 (0.421)		2147474	200.000	195.79
13	Trichlorofluoromethane	101	2.296	2.296 (0.457)		3119390	200.000	199.42
15	Acrolein	56	2.603	2.603 (0.518)		2896520	2000.00	1983.9
16	Acetone	43	2.722	2.722 (0.541)		1907733	400.000	400.17(A)
17	1,1-Dichloroethene	96	2.710	2.710 (0.539)		2115099	200.000	202.35(A)
18	Freon-113	151	2.722	2.722 (0.541)		1494012	200.000	200.47(A)

Data File: \\qcanoh04\dd\chem\MSV\3aux11.i\J40914B-1C.D \UAN430100
 Report Date: 15-Sep-2004 12:45

Compounds	QUANT SIG	MASS	AMOUNTS					
			RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)
		====	==	=====	=====	=====	=====	=====
		142	2.828	2.828 (0.562)		3111800	200.000	201.42 (A)
Iodomethane		76	2.899	2.899 (0.577)		7890340	200.000	202.08 (A)
Carbon Disulfide		84	3.077	3.077 (0.612)		2622600	200.000	200.02 (A)
Methylene Chloride		41	2.935	2.935 (0.584)		2423653	2000.00	1747.2
Acetonitrile		53	3.254	3.254 (0.647)		8435321	2000.00	1955.4
Acrylonitrile		73	3.301	3.301 (0.657)		6861860	200.000	204.77 (A)
Methyl tert-butyl ether		96	3.301	3.301 (0.657)		2403695	200.000	193.85
trans-1,2-Dichloroethene		86	3.526	3.526 (0.701)		460062	200.000	200.35 (A)
Hexane		43	3.656	3.656 (0.727)		4234296	200.000	213.40 (A)
Vinyl acetate		63	3.633	3.633 (0.722)		4472069	200.000	197.70
1,1-Dichloroethane		59	3.148	3.148 (0.626)		3293397	4000.00	3622.6 (A)
tert-Butyl Alcohol		43	4.082	4.082 (0.812)		2412455	400.000	376.83 (A)
2-Butanone		96				4951179	400.000	389.38
M 1,2-Dichloroethene (total)		96	4.094	4.094 (0.814)		2547484	200.000	195.53
cis-1,2-dichloroethene		77	4.094	4.094 (0.814)		2760188	200.000	202.48 (A)
2,2-Dichloropropane		128	4.284	4.284 (0.852)		1158230	200.000	191.98
Bromochloromethane		83	4.343	4.343 (0.864)		4417574	200.000	194.27
Chloroform		42	4.331	4.331 (0.861)		658534	200.000	199.60
Tetrahydrofuran		97	4.508	4.508 (0.896)		3459032	200.000	201.42 (A)
1,1,1-Trichloroethane		75	4.639	4.639 (0.922)		3312291	200.000	206.94 (A)
Dichloropropene		117	4.650	4.650 (0.925)		2829152	200.000	209.81 (A)
Carbon Tetrachloride		62	4.804	4.804 (0.955)		3659770	200.000	196.98
1,2-Dichloroethane		78	4.816	4.816 (0.958)		10307958	200.000	191.55
Benzene		130	5.337	5.337 (1.061)		2454964	200.000	199.89
Trichloroethene		63	5.514	5.514 (1.096)		2581062	200.000	194.44
1,2-Dichloropropane		88	5.621	5.621 (1.118)		961094	10000.0	8446.6 (A)
1,4-Dioxane		93	5.621	5.621 (1.118)		1453325	200.000	197.58
Dibromomethane		83	5.739	5.739 (1.141)		3494837	200.000	200.14 (A)
Bromodichloromethane		63	5.988	5.988 (1.191)		3196023	400.000	428.19 (A)
2-Chloroethyl vinyl ether		75	6.130	6.130 (1.219)		4425099	200.000	207.11 (A)
cis-1,3-Dichloropropene		43	6.248	6.248 (1.242)		4708742	400.000	409.86 (A)
4-Methyl-2-pentanone		91	6.425	6.425 (0.838)		11034583	200.000	200.23 (A)
Toluene		75	6.603	6.603 (0.861)		4148070	200.000	209.93 (A)
trans-1,3-Dichloropropene		69	6.674	6.674 (0.870)		3684757	200.000	216.74 (A)
Ethyl Methacrylate		97	6.769	6.769 (0.883)		2198790	200.000	196.70
1,1,2-Trichloroethane		76	6.922	6.922 (0.903)		4169551	200.000	197.96
1,3-Dichloropropane		164	6.934	6.934 (0.904)		1776877	200.000	199.01
Tetrachloroethene		43	6.982	6.982 (0.911)		3736062	400.000	418.68 (A)
2-Hexanone		129	7.135	7.135 (0.931)		2430585	200.000	203.85 (A)
Dibromochloromethane		107	7.242	7.242 (0.944)		2251324	200.000	204.40 (A)
1,2-Dibromoethane		112	7.703	7.703 (1.005)		7055771	200.000	197.73
Chlorobenzene		131	7.774	7.774 (1.014)		2491620	200.000	200.99 (A)
1,1,1,2-Tetrachloroethane		106	7.798	7.798 (1.017)		3790824	200.000	210.37 (A)
Ethylbenzene		106	7.904	7.904 (1.031)		9470698	400.000	408.06 (A)
m + p-Xylene		106				14172021	600.000	614.39
M 63 Xylenes (total)		106	8.283	8.283 (1.080)		4701323	200.000	206.33 (A)
Xylene-o		104	8.295	8.295 (1.082)		8616448	200.000	211.47 (A)
Styrene								

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng) ON-COL (ng)
66 Bromoform	====	173	8.472	8.472 (1.105)	1716386	200.000	211.78 (A)
67 Isopropylbenzene	105	8.626	8.626 (1.125)	10843610	200.000	213.68 (A)	
68 1,1,2,2-Tetrachloroethane	83	8.898	8.898 (0.898)	3053504	200.000	196.70	
69 1,4-Dichloro-2-butene	53	8.958	8.958 (0.904)	1088280	200.000	213.07 (A)	
70 1,2,3-Trichloropropane	110	8.946	8.946 (0.903)	994195	200.000	195.17	
71 Bromobenzene	156	8.922	8.922 (0.901)	3020894	200.000	196.61	
72 n-Propylbenzene	120	9.029	9.029 (0.912)	2967729	200.000	207.17 (A)	
73 2-Chlorotoluene	126	9.111	9.111 (0.920)	2837950	200.000	200.05 (A)	
74 1,3,5-Trimethylbenzene	105	9.194	9.194 (0.928)	10066915	200.000	211.19 (A)	
75 4-Chlorotoluene	126	9.218	9.218 (0.931)	2975348	200.000	196.33	
76 tert-Butylbenzene	119	9.514	9.514 (0.961)	7991718	200.000	206.94 (A)	
77 1,2,4-Trimethylbenzene	105	9.561	9.561 (0.965)	10631151	200.000	208.92 (A)	
78 sec-Butylbenzene	105	9.727	9.727 (0.982)	10794030	200.000	206.50 (A)	
79 4-Isopropyltoluene	119	9.869	9.869 (0.996)	9271210	200.000	213.18 (A)	
80 1,3-Dichlorobenzene	146	9.845	9.845 (0.994)	5493140	200.000	193.66	
81 1,4-Dichlorobenzene	146	9.928	9.928 (1.002)	5803120	200.000	195.54	
82 n-Butylbenzene	91	10.271	10.271 (1.037)	8332031	200.000	214.47 (A)	
83 1,2-Dichlorobenzene	146	10.295	10.295 (1.039)	5384330	200.000	192.72	
84 1,2-Dibromo-3-chloropropane	157	11.052	11.052 (1.116)	528598	200.000	201.08 (A)	
85 1,2,4-Trichlorobenzene	180	11.892	11.892 (1.201)	2505741	200.000	210.67 (A)	
86 Hexachlorobutadiene	225	12.070	12.070 (1.219)	979903	200.000	200.36 (A)	
87 Naphthalene	128	12.129	12.129 (1.225)	6432622	200.000	229.14 (A)	
88 1,2,3-Trichlorobenzene	180	12.377	12.377 (1.250)	1787787	200.000	209.60 (A)	
98 Cyclohexane	56	4.568	4.568 (0.908)	3724145	200.000	216.58 (A)	
143 Methyl Acetate	43	2.982	2.982 (0.593)	3693009	400.000	384.10	
144 Methylcyclohexane	83	5.514	5.514 (1.096)	3054148	200.000	211.66	
141 1,3,5-Trichlorobenzene	180	11.277	11.277 (1.139)	3087088	200.000	196.62	

QC Flag Legend

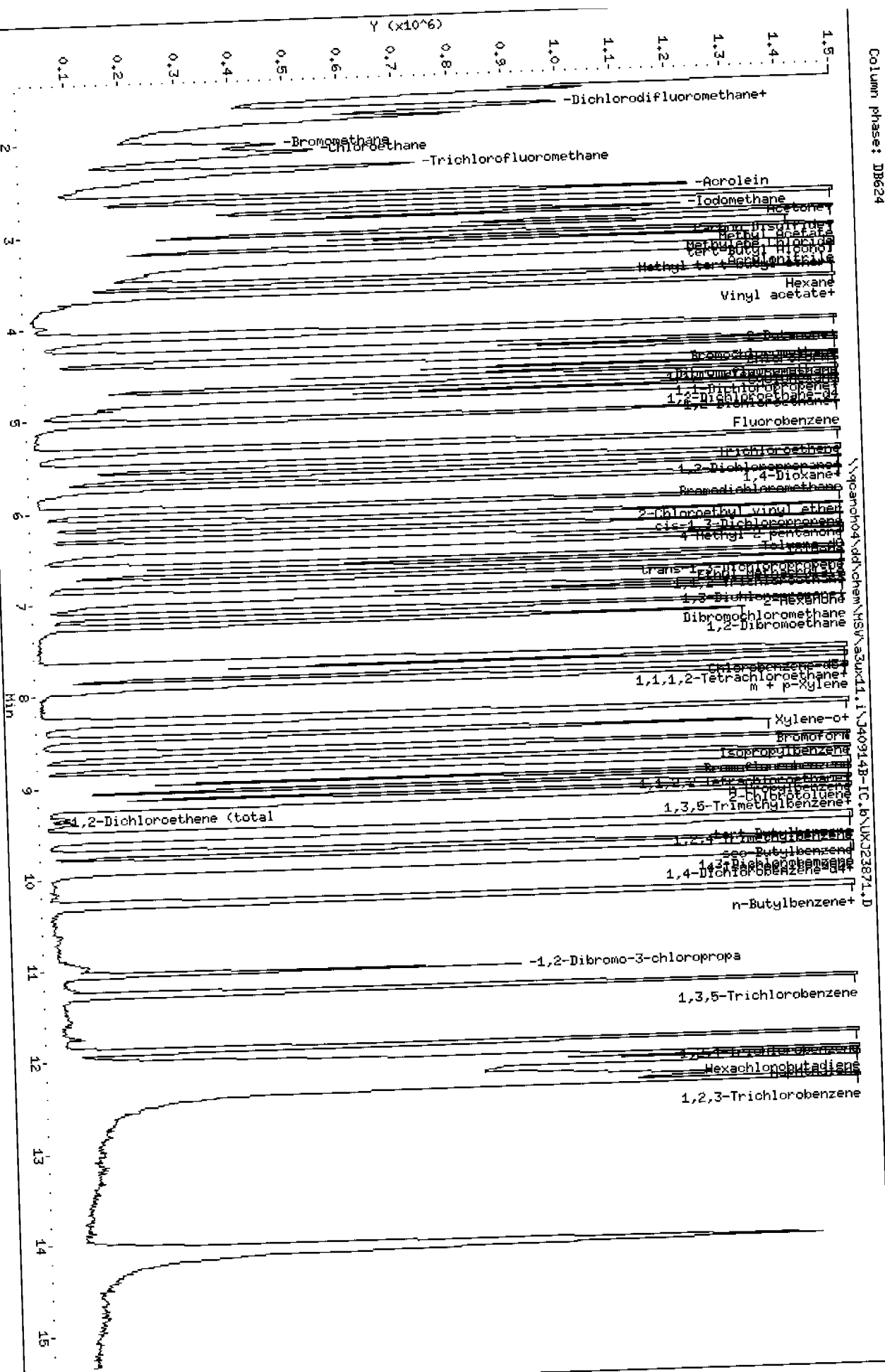
A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\qcancn04\dd\chem\HSV\z3ux11.i\J40914B-IC.b\JKJ23871.D
 Date : 14-SEP-2004 14:10
 Client ID:
 Sample Info: 100NG-IC
 Purge Volume: 5.0
 Column Phase: DB624

Instrument: z3ux11.i

Operator: 43582

Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.D\UXJ23871.D
Report Date: 15-Sep-2004 12:45

STL North Canton

VOLATILE REPORT SW-846 Method
Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\UXJ23871.D

Lab Smp Id: 100NG-IC
Inj Date : 14-SEP-2004 14:10 Inst ID: A3UX11.i
Operator : 43582
Smp Info : 100NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 5
Comment : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40914B-IC.b\8260LLUX11.m Quant Type: ISTD
Meth Date : 15-Sep-2004 12:45 evansl Cal File: UXJ23875.D
Cal Date : 14-SEP-2004 15:41 Calibration Sample, Level: 5
Als bottle: 3
Dil Factor: 1.00000 Compound Sublist: 2-8260.SUB
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/V_o

Name	Value	Description
DF	1.000	Dilution Factor
V _o	5.000	Sample volume

Compounds	QUANT SIG	MASS	RT	EXP RT	REL RT	RESPONSE	AMOUNTS	
							CAL-AMT (ng)	ON-COL (ng)
*		96	5.029	5.029 (1.000)		2294078	50.0000	
*		117	7.680	7.680 (1.000)		1821500	50.0000	
*		152	9.904	9.904 (1.000)		999916	50.0000	
\$		113	4.473	4.473 (0.889)		1074908	100.000	100.64
\$		65	4.745	4.745 (0.944)		1555269	100.000	105.89
\$		98	6.378	6.378 (0.831)		4394161	100.000	100.42
\$		95	8.780	8.780 (1.143)		1899831	100.000	102.18
\$		85	1.527	1.527 (0.304)		1124779	100.000	93.783
8		50	1.680	1.680 (0.334)		1934079	100.000	90.319
9		62	1.763	1.763 (0.351)		1367463	100.000	93.356
10		94	2.047	2.047 (0.407)		634017	100.000	91.256
11		64	2.130	2.130 (0.424)		1013498	100.000	94.147
12		101	2.296	2.296 (0.457)		1404082	100.000	91.454
13		56	2.615	2.615 (0.520)		1447814	1000.00	1010.3
15		43	2.722	2.722 (0.541)		973885	200.000	198.82
16		96	2.710	2.710 (0.539)		950397	100.000	92.638
17		151	2.734	2.734 (0.544)		619208	100.000	93.888
18								

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
19 Iodomethane	142	2.828	2.828 (0.562)	1515000	100.000	99.909	
20 Carbon Disulfide	76	2.899	2.899 (0.577)	3602731	100.000	94.011	
21 Methylene Chloride	84	3.077	3.077 (0.612)	1333682	100.000	99.833	
22 Acetonitrile	41	2.935	2.935 (0.584)	1365511	1000.00	1002.9	
23 Acrylonitrile	53	3.254	3.254 (0.647)	4234501	1000.00	1000.1	
24 Methyl tert-butyl ether	73	3.302	3.302 (0.657)	3440137	100.000	104.59	
25 trans-1,2-Dichloroethene	96	3.302	3.302 (0.657)	1175346	100.000	96.574	
26 Hexane	86	3.526	3.526 (0.701)	195007	100.000	95.842	
27 Vinyl acetate	43	3.657	3.657 (0.727)	2062282	100.000	105.89	
28 1,1-Dichloroethane	63	3.633	3.633 (0.722)	2173037	100.000	97.878	
29 tert-Butyl Alcohol	59	3.148	3.148 (0.626)	1816490	2000.00	2035.8 (A)	
30 2-Butanone	43	4.083	4.083 (0.812)	1242482	200.000	197.74	
M 31 1,2-Dichloroethene (total)	96			2422890	200.000	194.13	
32 cis-1,2-dichloroethene	96	4.094	4.094 (0.814)	1247544	100.000	97.561	
33 2,2-Dichloropropane	77	4.106	4.106 (0.816)	1285494	100.000	96.081	
34 Bromochloromethane	128	4.284	4.284 (0.852)	577210	100.000	97.480	
35 Chloroform	83	4.343	4.343 (0.864)	2173231	100.000	97.374	
36 Tetrahydrofuran	42	4.331	4.331 (0.861)	344585	100.000	101.91	
37 1,1,1-Trichloroethane	97	4.509	4.509 (0.896)	1598802	100.000	94.853	
38 1,1-Dichloropropene	75	4.639	4.639 (0.922)	1510060	100.000	96.123	
39 Carbon Tetrachloride	117	4.651	4.651 (0.925)	1268602	100.000	95.854	
40 1,2-Dichloroethane	62	4.816	4.816 (0.958)	1797145	100.000	98.552	
41 Benzene	78	4.816	4.816 (0.958)	5026234	100.000	95.163	
42 Trichloroethene	130	5.337	5.337 (1.061)	1185674	100.000	98.362	
43 1,2-Dichloropropane	63	5.526	5.526 (1.099)	1270520	100.000	97.515	
44 1,4-Dioxane	88	5.621	5.621 (1.118)	597256	5000.00	5347.9 (A)	
45 Dibromomethane	93	5.621	5.621 (1.118)	727774	100.000	100.80	
46 Bromodichloromethane	83	5.751	5.751 (1.144)	1694447	100.000	98.863	
47 2-Chloroethyl vinyl ether	63	5.988	5.988 (1.191)	1586020	200.000	216.50 (A)	
48 cis-1,3-Dichloropropene	75	6.130	6.130 (1.219)	2139621	100.000	102.03	
49 4-Methyl-2-pentanone	43	6.248	6.248 (1.242)	2343412	200.000	207.82 (A)	
50 Toluene	91	6.437	6.437 (0.838)	5341010	100.000	98.181	
51 trans-1,3-Dichloropropene	75	6.603	6.603 (0.860)	2025722	100.000	103.86	
52 Ethyl Methacrylate	69	6.674	6.674 (0.869)	1801585	100.000	107.35	
53 1,1,2-Trichloroethane	97	6.769	6.769 (0.881)	1103061	100.000	99.965	
54 1,3-Dichloropropane	76	6.922	6.922 (0.901)	2083135	100.000	100.19	
55 Tetrachloroethene	164	6.934	6.934 (0.903)	827625	100.000	93.905	
56 2-Hexanone	43	6.982	6.982 (0.909)	1792800	200.000	203.53 (A)	
57 Dibromochloromethane	129	7.135	7.135 (0.929)	1188499	100.000	100.98	
58 1,2-Dibromoethane	107	7.242	7.242 (0.943)	1119552	100.000	102.98	
59 Chlorobenzene	112	7.703	7.703 (1.003)	3458216	100.000	98.180	
60 1,1,1,2-Tetrachloroethane	131	7.774	7.774 (1.012)	1239765	100.000	101.31	
61 Ethylbenzene	106	7.798	7.798 (1.015)	1782706	100.000	100.22	
62 m + p-Xylene	106	7.905	7.905 (1.029)	4519264	200.000	197.26	
M 63 xylenes (total)	106			6782922	300.000	297.91	
64 Xylene-o	106	8.283	8.283 (1.079)	2263658	100.000	100.65	
65 Styrene	104	8.295	8.295 (1.080)	4147114	100.000	103.11	

Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\UXJ23871.D
 Report Date: 15-Sep-2004 12:45

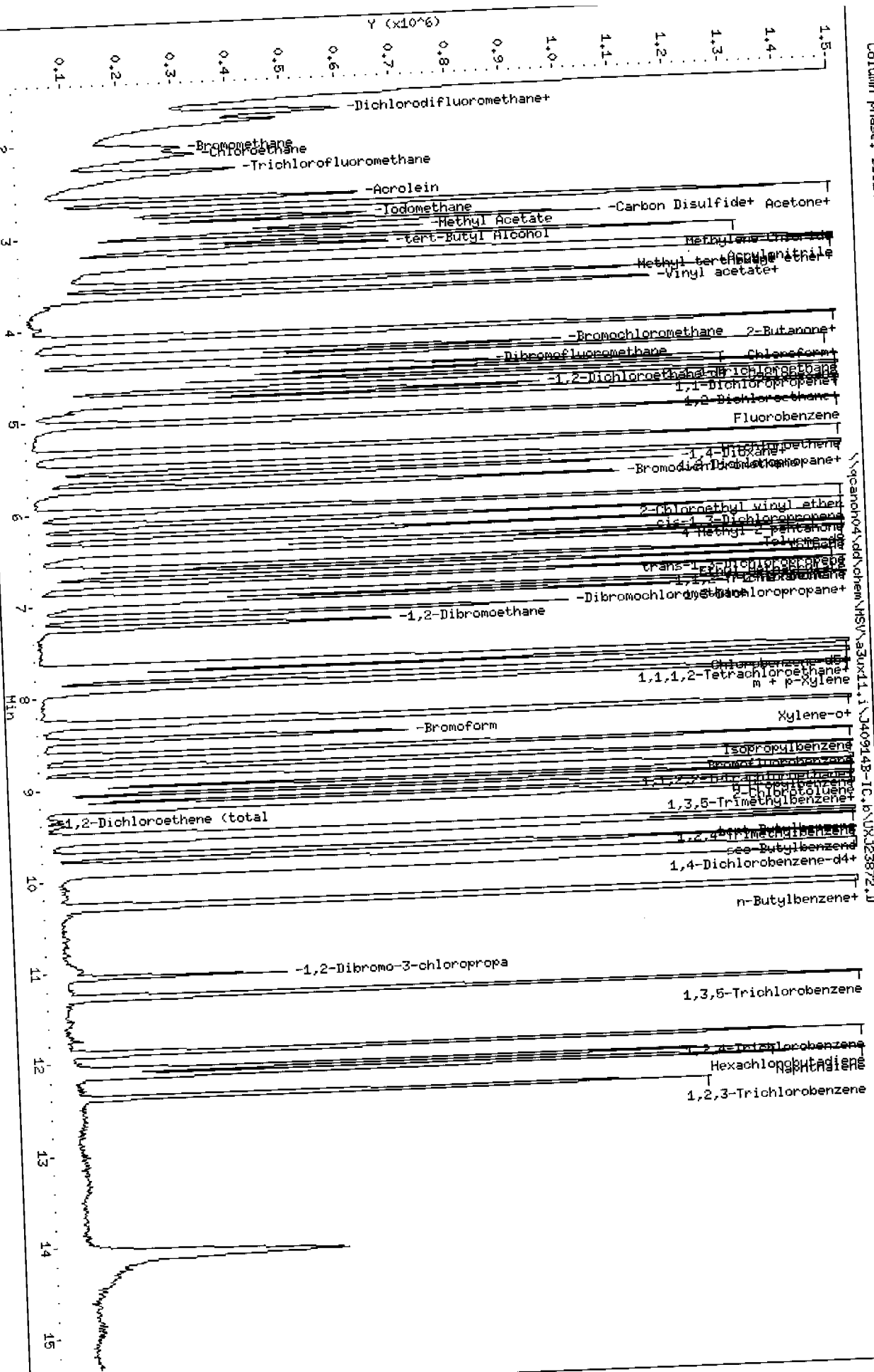
Compounds	QUANT SIG	MASS	AMOUNTS					
			RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)
66 Bromoform		173	8.473	8.473 (1.103)		820790	100.000	102.60
67 Isopropylbenzene		105	8.626	8.626 (1.123)		4943240	100.000	98.682
68 1,1,2,2-Tetrachloroethane		83	8.899	8.899 (0.898)		1522185	100.000	100.99
69 1,4-Dichloro-2-butene		53	8.958	8.958 (0.904)		523620	100.000	105.58
70 1,2,3-Trichloropropane		110	8.946	8.946 (0.903)		495410	100.000	100.16
71 Bromobenzene		156	8.934	8.934 (0.902)		1490126	100.000	99.884
72 n-Propylbenzene		120	9.029	9.029 (0.912)		1351668	100.000	97.179
73 2-Chlorotoluene		126	9.112	9.112 (0.920)		1351055	100.000	98.084
74 1,3,5-Trimethylbenzene		105	9.194	9.194 (0.928)		4630795	100.000	100.05
75 4-Chlorotoluene		126	9.218	9.218 (0.931)		1433771	100.000	97.438
76 tert-Butylbenzene		119	9.514	9.514 (0.961)		3670015	100.000	97.876
77 1,2,4-Trimethylbenzene		105	9.561	9.561 (0.965)		5001495	100.000	101.23
78 sec-Butylbenzene		105	9.727	9.727 (0.982)		4894571	100.000	96.437
79 4-Isopropyltoluene		119	9.869	9.869 (0.996)		4211412	100.000	99.734
80 1,3-Dichlorobenzene		146	9.845	9.845 (0.994)		2672507	100.000	97.033
81 1,4-Dichlorobenzene		146	9.928	9.928 (1.002)		2817287	100.000	97.770
82 n-Butylbenzene		91	10.271	10.271 (1.037)		3687678	100.000	97.759
83 1,2-Dichlorobenzene		146	10.295	10.295 (1.039)		2647056	100.000	97.579
84 1,2-Dibromo-3-chloropropane		157	11.052	11.052 (1.116)		261375	100.000	102.40
85 1,2,4-Trichlorobenzene		180	11.892	11.892 (1.201)		1158323	100.000	100.30
86 Hexachlorobutadiene		225	12.070	12.070 (1.219)		475139	100.000	97.147
87 Naphthalene		128	12.129	12.129 (1.225)		3076588	100.000	112.87
88 1,2,3-Trichlorobenzene		180	12.377	12.377 (1.250)		883388	100.000	106.66
98 Cyclohexane		56	4.568	4.568 (0.908)		1591353	100.000	94.291
143 Methyl Acetate		43	2.994	2.994 (0.595)		1827750	200.000	193.68
144 Methylcyclohexane		83	5.514	5.514 (1.096)		1298110	100.000	91.658
141 1,3,5-Trichlorobenzene		180	11.277	11.277 (1.139)		1469129	100.000	96.368

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Instrument: a30x11.i

Operator: 43582
 Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23872.D
Report Date: 15-Sep-2004 12:46

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23872.D
Lab Smp Id: 50NG-IC
Inj Date : 14-SEP-2004 14:33 Inst ID: a3ux11.i
Operator : 43582
Smp Info : 50NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 4
Comment : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m Quant Type: ISTD
Meth Date : 15-Sep-2004 12:46 evans1 Cal File: UXJ23875.D
Cal Date : 14-SEP-2004 15:41 Calibration Sample, Level: 4
Als bottle: 4
Dil Factor: 1.00000 Compound Sublist: 2-8260.SUB
Integrator: HP RTE
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/V_o

Name	Value	Description
DF	1.000	Dilution Factor
V _o	5.000	Sample volume

Compounds	QUANT SIG	MASS	AMOUNTS				
			RT	EXP RT	REL RT	RESPONSE	(ng) ON-COL (ng)
*	96	5.029	5.029 (1.000)	2329625	50.0000		
1 Fluorobenzene	117	7.668	7.668 (1.000)	1858993	50.0000		
2 Chlorobenzene-d5	152	9.904	9.904 (1.000)	990958	50.0000		
3 1,4-Dichlorobenzene-d4	113	4.473	4.473 (0.889)	558177	50.0000	51.462	
4 Dibromofluoromethane	65	4.745	4.745 (0.944)	733545	50.0000	49.183	
5 1,2-Dichloroethane-d4	98	6.378	6.378 (0.832)	2256469	50.0000	50.525	
6 Toluene-d8	95	8.780	8.780 (1.145)	943942	50.0000	49.744	
7 Bromofluorobenzene	85	1.527	1.527 (0.304)	586151	50.0000	48.127	
8 Dichlorodifluoromethane	50	1.680	1.680 (0.334)	1022123	50.0000	47.004	
9 Chloromethane	62	1.763	1.763 (0.351)	726372	50.0000	48.832	
10 Vinyl Chloride	94	2.047	2.047 (0.407)	352300	50.0000	49.934	
11 Bromomethane	64	2.130	2.130 (0.424)	545580	50.0000	49.907	
12 Chloroethane	101	2.296	2.296 (0.457)	757893	50.0000	48.612	
13 Trichlorofluoromethane	56	2.615	2.615 (0.520)	720539	500.000	495.15	
15 Acrolein	43	2.722	2.722 (0.541)	530389	100.000	102.36	
16 Acetone	96	2.710	2.710 (0.539)	541982	50.0000	52.022	
17 1,1-Dichloroethene	151	2.722	2.722 (0.541)	390502	50.0000	59.593	
18 Freon-113							

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.D\UAN4201404
 Report Date: 15-Sep-2004 12:46

Compounds	QUANT SIG	MASS	AMOUNTS					
			RT	EXP RT	REL RT	RESPONSE	CAL-AMT	ON-COL
							(ng)	(ng)
			=	=	=	=	=	=
			142	2.828	2.828 (0.562)	777257	50.0000	50.475
19 Iodomethane		76	2.899	2.899 (0.577)		1989456	50.0000	51.121
20 Carbon Disulfide		84	3.077	3.077 (0.612)		741159	50.0000	50.431
21 Methylene Chloride		41	2.935	2.935 (0.584)		667504	500.000	482.78
22 Acetonitrile		53	3.254	3.254 (0.647)		2117146	500.000	492.41
23 Acrylonitrile		73	3.302	3.302 (0.657)		1733984	50.0000	51.915
24 Methyl tert-butyl ether		96	3.302	3.302 (0.657)		623149	50.0000	50.421
25 trans-1,2-Dichloroethene		86	3.526	3.526 (0.701)		115096	50.0000	57.272
26 Hexane		43	3.657	3.657 (0.727)		979072	50.0000	49.506
27 Vinyl acetate		63	3.633	3.633 (0.722)		1126418	50.0000	49.962
28 1,1-Dichloroethane		59	3.148	3.148 (0.626)		896352	1000.00	989.22
29 tert-Butyl Alcohol		43	4.083	4.083 (0.812)		623229	100.000	97.672
30 2-Butanone		96				1262694	100.000	99.671
M 31 1,2-Dichloroethene (total)		96	4.094	4.094 (0.814)		639545	50.0000	49.251
32 cis-1,2-dichloroethene		77	4.106	4.106 (0.816)		684525	50.0000	50.382
33 2,2-Dichloropropane		128	4.284	4.284 (0.852)		299703	50.0000	49.842
34 Bromochloromethane		83	4.343	4.343 (0.864)		1133371	50.0000	50.007
35 Chloroform		42	4.331	4.331 (0.861)		170087	50.0000	48.338
36 Tetrahydrofuran		97	4.508	4.508 (0.896)		892252	50.0000	52.127
37 1,1,1-Trichloroethane		75	4.639	4.639 (0.922)		833818	50.0000	52.267
38 1,1-Dichloropropene		117	4.650	4.650 (0.925)		724043	50.0000	49.362
39 Carbon Tetrachloride		62	4.816	4.816 (0.958)		914079	50.0000	49.261
40 1,2-Dichloroethane		78	4.816	4.816 (0.958)		2642144	50.0000	50.875
41 Benzene		130	5.337	5.337 (1.061)		622755	50.0000	49.474
42 Trichloroethene		63	5.526	5.526 (1.099)		654580	50.0000	2681.8 (A)
43 1,2-Dichloropropane		88	5.621	5.621 (1.118)		304150	2500.00	50.275
44 1,4-Dioxane		93	5.621	5.621 (1.118)		368591	50.0000	49.580
45 Dibromomethane		83	5.751	5.751 (1.144)		862931	50.0000	104.55
46 Bromodichloromethane		63	5.988	5.988 (1.191)		777773	100.000	49.761
47 2-Chloroethyl vinyl ether		75	6.130	6.130 (1.219)		1059681	50.0000	102.59
48 cis-1,3-Dichloropropene		43	6.248	6.248 (1.242)		1174753	100.000	49.955
49 4-Methyl-2-pentanone		91	6.437	6.437 (0.840)		2773477	50.0000	50.651
50 Toluene		75	6.603	6.603 (0.861)		1008249	50.0000	52.165
51 trans-1,3-Dichloropropene		69	6.674	6.674 (0.870)		893443	50.0000	50.102
52 Ethyl Methacrylate		97	6.769	6.769 (0.883)		564234	50.0000	49.040
53 1,1,2-Trichloroethane		76	6.922	6.922 (0.903)		1040578	50.0000	50.654
54 1,3-Dichloropropane		164	6.934	6.934 (0.904)		455621	50.0000	102.93
55 Tetrachloroethene		43	6.982	6.982 (0.911)		925357	100.000	49.450
56 2-Hexanone		129	7.135	7.135 (0.931)		593974	50.0000	50.647
57 Dibromochloromethane		107	7.242	7.242 (0.944)		561972	50.0000	48.975
58 1,2-Dibromoethane		112	7.703	7.703 (1.005)		1760577	50.0000	49.009
59 Chlorobenzene		131	7.774	7.774 (1.014)		612067	50.0000	50.081
60 1,1,1,2-Tetrachloroethane		106	7.798	7.798 (1.017)		909162	50.0000	101.65
61 Ethylbenzene		106	7.905	7.905 (1.031)		2376751	100.000	151.84
62 m + p-Xylene		106				3528868	150.000	50.192
M 63 Xylenes (total)		106	8.283	8.283 (1.080)		1152117	50.0000	50.479
64 Xylene-o		106	8.295	8.295 (1.082)		2072070	50.0000	
65 Styrene		104						

Compounds	QUANT SIG	MASS	RT	EXP RT	REL RT	RESPONSE	AMOUNTS	
							CAL-AMT (ng)	ON-COL (ng)
66 Bromoform	====	173	8.473	8.473 (1.105)		406002	50.0000	49.727
67 Isopropylbenzene	====	105	8.626	8.626 (1.125)		2624726	50.0000	51.341
68 1,1,2,2-Tetrachloroethane	====	83	8.899	8.899 (0.898)		750188	50.0000	50.220
69 1,4-Dichloro-2-butene	====	53	8.958	8.958 (0.904)		2494440	50.0000	50.752
70 1,2,3-Trichloropropane	====	110	8.946	8.946 (0.903)		246053	50.0000	50.196
71 Bromobenzene	====	156	8.934	8.934 (0.902)		745059	50.0000	50.393
72 n-Propylbenzene	====	120	9.029	9.029 (0.912)		716364	50.0000	51.969
73 2-Chlorotoluene	====	126	9.111	9.111 (0.920)		694404	50.0000	50.868
74 1,3,5-Trimethylbenzene	====	105	9.194	9.194 (0.928)		2372465	50.0000	51.722
75 4-Chlorotoluene	====	126	9.218	9.218 (0.931)		738082	50.0000	50.613
76 tert-Butylbenzene	====	119	9.514	9.514 (0.961)		1923678	50.0000	51.766
77 1,2,4-Trimethylbenzene	====	105	9.561	9.561 (0.965)		2551790	50.0000	52.114
78 sec-Butylbenzene	====	105	9.727	9.727 (0.982)		2633655	50.0000	52.360
79 4-Isopropyltoluene	====	119	9.869	9.869 (0.996)		2250860	50.0000	53.786
80 1,3-Dichlorobenzene	====	146	9.845	9.845 (0.994)		1354905	50.0000	49.638
81 1,4-Dichlorobenzene	====	146	9.928	9.928 (1.002)		1437001	50.0000	50.320
82 n-Butylbenzene	====	91	10.271	10.271 (1.037)		2001622	50.0000	53.542
83 1,2-Dichlorobenzene	====	146	10.295	10.295 (1.039)		1339377	50.0000	49.820
84 1,2-Dibromo-3-chloropropane	====	157	11.052	11.052 (1.116)		127154	50.0000	50.267
85 1,2,4-Trichlorobenzene	====	180	11.892	11.892 (1.201)		576092	50.0000	50.333
86 Hexachlorobutadiene	====	225	12.070	12.070 (1.219)		272701	50.0000	54.568
87 Naphthalene	====	128	12.129	12.129 (1.225)		1382075	50.0000	51.162
88 1,2,3-Trichlorobenzene	====	180	12.377	12.377 (1.250)		415515	50.0000	50.625
98 Cyclohexane	====	56	4.568	4.568 (0.908)		952028	50.0000	55.549
143 Methyl Acetate	====	43	2.994	2.994 (0.595)		911479	100.000	95.114
144 Methylcyclohexane	====	83	5.514	5.514 (1.096)		793891	50.0000	55.200
141 1,3,5-Trichlorobenzene	====	180	11.277	11.277 (1.139)		752113	50.0000	49.781

QC Flag Legend

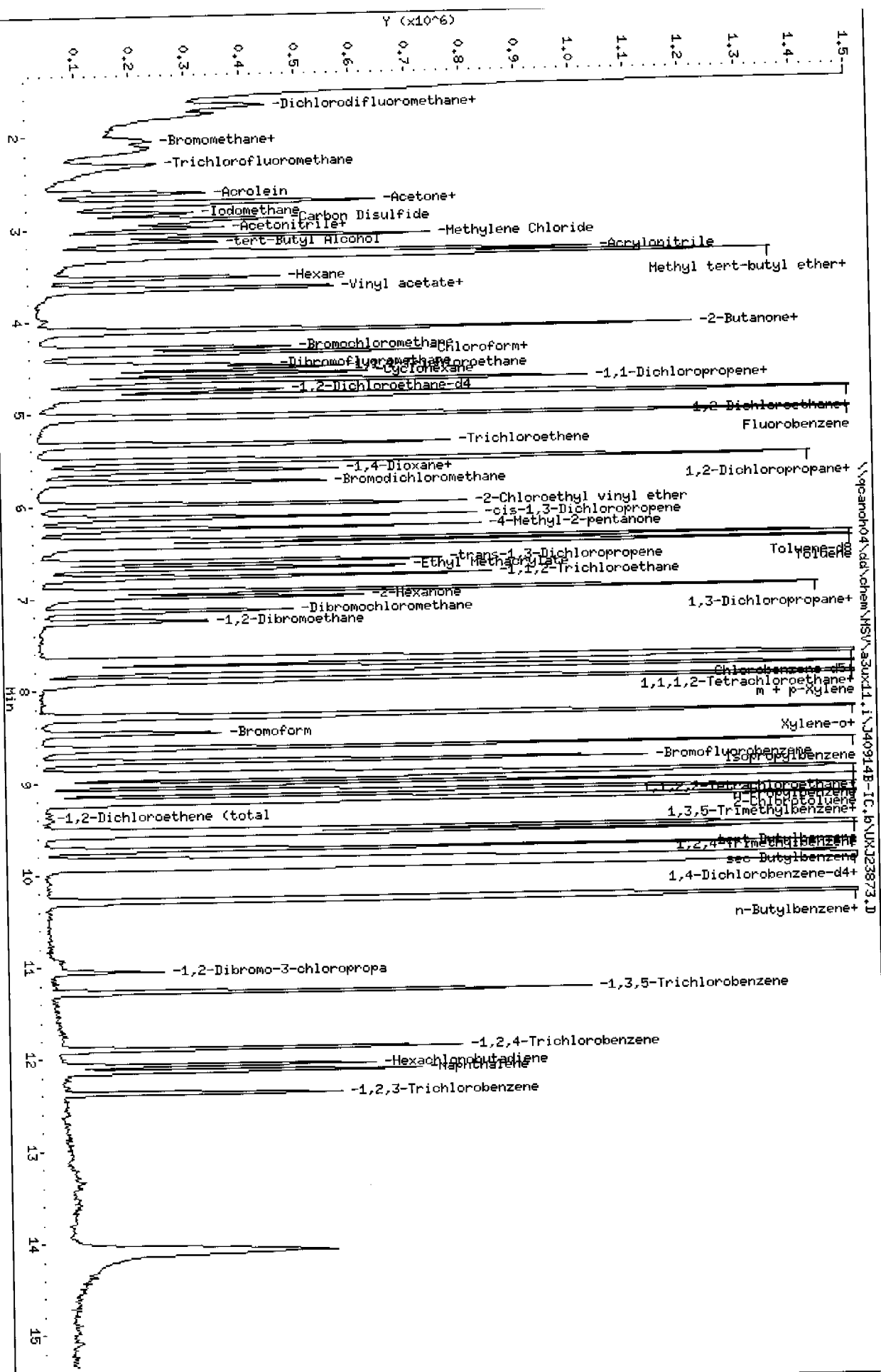
A - Target compound detected but, quantitated amount exceeded maximum amount.

Client ID:
 Sample Info: 25NG-IC
 Purge Volume: 5.0

Column Phase: DB624

Instrument: a30x11.i

Operator: 43582
 Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23873.D
Report Date: 15-Sep-2004 12:47

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23873.D
Lab Smp Id: 25NG-IC
Inj Date : 14-SEP-2004 14:57
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 25NG-IC
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,1,3
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:46 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 5 Calibration Sample, Level: 3
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ng)
*	1 Fluorobenzene	96	5.029	5.029 (1.000)	2302064	50.0000		
*	2 Chlorobenzene-d5	117	7.668	7.668 (1.000)	1820780	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	1002006	50.0000		
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.889)	263579	25.0000	24.592	
\$	5 1,2-Dichloroethane-d4	65	4.745	4.745 (0.944)	360514	25.0000	24.461	
\$	6 Toluene-d8	98	6.378	6.378 (0.832)	1137706	25.0000	26.009	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.145)	468289	25.0000	25.196	
8	Dichlorodifluoromethane	85	1.526	1.526 (0.304)	324283	25.0000	26.945	
9	Chloromethane	50	1.680	1.680 (0.334)	523520	25.0000	24.363	
10	Vinyl Chloride	62	1.775	1.775 (0.353)	382674	25.0000	26.034	
11	Bromomethane	94	2.047	2.047 (0.407)	163334	25.0000	23.428	
12	Chloroethane	64	2.130	2.130 (0.424)	280413	25.0000	25.958	
13	Trichlorofluoromethane	101	2.296	2.296 (0.457)	399606	25.0000	25.938	
15	Acrolein	56	2.615	2.615 (0.520)	360127	250.000	250.44	
16	Acetone	43	2.722	2.722 (0.541)	262942	50.0000	48.200	
17	1,1-Dichloroethene	96	2.710	2.710 (0.539)	229496	25.0000	22.292	
18	Freon-113	151	2.722	2.722 (0.541)	126246	25.0000	19.380	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXU45813.D
 Report Date: 15-Sep-2004 12:47

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT	ON-COL
							(ng)	(ng)
Iodomethane		142	2.828	2.828 (0.562)		375596	25.0000	24.683
20 Carbon Disulfide		76	2.899	2.899 (0.577)		854870	25.0000	22.230
21 Methylene Chloride		84	3.077	3.077 (0.612)		413458	25.0000	24.263
22 Acetonitrile		41	2.946	2.946 (0.586)		352454	250.000	257.97
23 Acrylonitrile		53	3.254	3.254 (0.647)		1072529	250.000	252.44
24 Methyl tert-butyl ether		73	3.301	3.301 (0.657)		855584	25.0000	25.923
25 trans-1,2-Dichloroethene		96	3.313	3.313 (0.659)		291530	25.0000	23.871
26 Hexane		86	3.526	3.526 (0.701)		38256	25.0000	19.484
27 Vinyl acetate		43	3.656	3.656 (0.727)		490495	25.0000	25.099
28 1,1-Dichloroethane		63	3.633	3.633 (0.722)		530330	25.0000	23.804
29 tert-Butyl Alcohol		59	3.148	3.148 (0.626)		445116	500.000	497.11
30 2-Butanone		43	4.082	4.082 (0.812)		314848	50.0000	49.934
M 31 1,2-Dichloroethene (total)		96				603009	50.0000	48.145
32 cis-1,2-dichloroethene		96	4.094	4.094 (0.814)		311479	25.0000	24.274
33 2,2-Dichloroproppane		77	4.106	4.106 (0.816)		318520	25.0000	23.724
34 Bromochloromethane		128	4.284	4.284 (0.852)		145392	25.0000	24.469
35 Chloroform		83	4.343	4.343 (0.864)		541044	25.0000	24.158
36 Tetrahydrofuran		42	4.331	4.331 (0.861)		78777	25.0000	22.465
37 1,1,1-Trichloroethane		97	4.508	4.508 (0.896)		386062	25.0000	22.825
38 1,1-Dichloropropene		75	4.639	4.639 (0.922)		352859	25.0000	22.383
39 Carbon Tetrachloride		117	4.650	4.650 (0.925)		282428	25.0000	21.266
40 1,2-Dichloroethane		62	4.816	4.816 (0.958)		443026	25.0000	24.210
41 Benzene		78	4.816	4.816 (0.958)		1294581	25.0000	24.426
42 Trichloroethene		130	5.337	5.337 (1.061)		289238	25.0000	23.912
43 1,2-Dichloropropane		63	5.526	5.526 (1.099)		313009	25.0000	23.941
44 1,4-Dioxane		88	5.621	5.621 (1.118)		148470	1250.00	1324.8 (A)
45 Dibromomethane		93	5.621	5.621 (1.118)		175336	25.0000	24.202
46 Bromodichloromethane		83	5.751	5.751 (1.144)		415519	25.0000	24.160
47 2-Chloroethyl vinyl ether		63	5.987	5.987 (1.191)		367975	50.0000	50.055
48 cis-1,3-Dichloropropene		75	6.129	6.129 (1.219)		502234	25.0000	23.866
49 4-Methyl-2-pentanone		43	6.248	6.248 (1.242)		557786	50.0000	49.294
50 Toluene		91	6.437	6.437 (0.840)		1338715	25.0000	24.618
51 trans-1,3-Dichloropropene		75	6.603	6.603 (0.861)		468202	25.0000	24.015
52 Ethyl Methacrylate		69	6.674	6.674 (0.870)		409677	25.0000	24.422
53 1,1,2-Trichloroethane		97	6.768	6.768 (0.883)		270822	25.0000	24.553
54 1,3-Dichloropropane		76	6.922	6.922 (0.903)		513387	25.0000	24.702
55 Tetrachloroethene		164	6.934	6.934 (0.904)		201576	25.0000	22.880
56 2-Hexanone		43	6.981	6.981 (0.911)		441195	50.0000	50.107
57 Dibromochloromethane		129	7.135	7.135 (0.931)		291299	25.0000	24.760
58 1,2-Dibromoethane		107	7.254	7.254 (0.946)		264386	25.0000	24.328
59 Chlorobenzene		112	7.703	7.703 (1.005)		859376	25.0000	24.408
60 1,1,1,2-Tetrachloroethane		131	7.774	7.774 (1.014)		297129	25.0000	24.291
61 Ethylbenzene		106	7.798	7.798 (1.017)		427629	25.0000	24.050
62 m + p-Xylene		106	7.904	7.904 (1.031)		1125922	50.0000	49.164
M 63 Xylenes (total)		106	8.283	8.283 (1.080)		1692886	75.0000	74.383
64 Xylene-o		106	8.295	8.295 (1.082)		566964	25.0000	25.218
65 Styrene		104				985947	25.0000	24.523

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)
66 Bromoform	====	173	8.472	8.472 (1.105)		195832	25.0000	24.489
67 Isopropylbenzene		105	8.626	8.626 (1.125)		1187991	25.0000	23.725
68 1,1,2,2-Tetrachloroethane		83	8.898	8.898 (0.898)		370501	25.0000	24.529
69 1,4-Dichloro-2-butene		53	8.958	8.958 (0.904)		119797	25.0000	24.105
70 1,2,3-Trichloropropane		110	8.946	8.946 (0.903)		121026	25.0000	24.417
71 Bromobenzene		156	8.922	8.922 (0.901)		361498	25.0000	24.181
72 n-Propylbenzene		120	9.029	9.029 (0.912)		315155	25.0000	22.611
73 2-Chlorotoluene		126	9.111	9.111 (0.920)		330096	25.0000	23.914
74 1,3,5-Trimethylbenzene		105	9.194	9.194 (0.928)		1104880	25.0000	23.822
75 4-Chlorotoluene		126	9.218	9.218 (0.931)		353698	25.0000	23.987
76 tert-Butylbenzene		119	9.514	9.514 (0.961)		852078	25.0000	22.677
77 1,2,4-Trimethylbenzene		105	9.561	9.561 (0.965)		1171362	25.0000	23.658
78 sec-Butylbenzene		105	9.727	9.727 (0.982)		1145024	25.0000	22.513
79 4-Isopropyltoluene		119	9.869	9.869 (0.996)		954488	25.0000	22.557
80 1,3-Dichlorobenzene		146	9.845	9.845 (0.994)		654290	25.0000	23.706
81 1,4-Dichlorobenzene		146	9.928	9.928 (1.002)		690846	25.0000	23.925
82 n-Butylbenzene		91	10.271	10.271 (1.037)		825372	25.0000	21.835
83 1,2-Dichlorobenzene		146	10.295	10.295 (1.039)		651646	25.0000	23.972
84 1,2-Dibromo-3-chloropropane		157	11.052	11.052 (1.116)		63184	25.0000	24.703
85 1,2,4-Trichlorobenzene		180	11.892	11.892 (1.201)		261752	25.0000	22.617
86 Hexachlorobutadiene		225	12.070	12.070 (1.219)		126348	25.0000	23.115
87 Naphthalene		128	12.129	12.129 (1.225)		610547	25.0000	22.352
88 1,2,3-Trichlorobenzene		180	12.377	12.377 (1.250)		182675	25.0000	22.011
98 Cyclohexane		56	4.568	4.568 (0.908)		334819	25.0000	19.770
143 Methyl Acetate		43	2.994	2.994 (0.595)		462866	50.0000	48.879
144 Methylcyclohexane		83	5.514	5.514 (1.096)		275331	25.0000	19.373
141 1,3,5-Trichlorobenzene		180	11.277	11.277 (1.139)		338982	25.0000	22.189

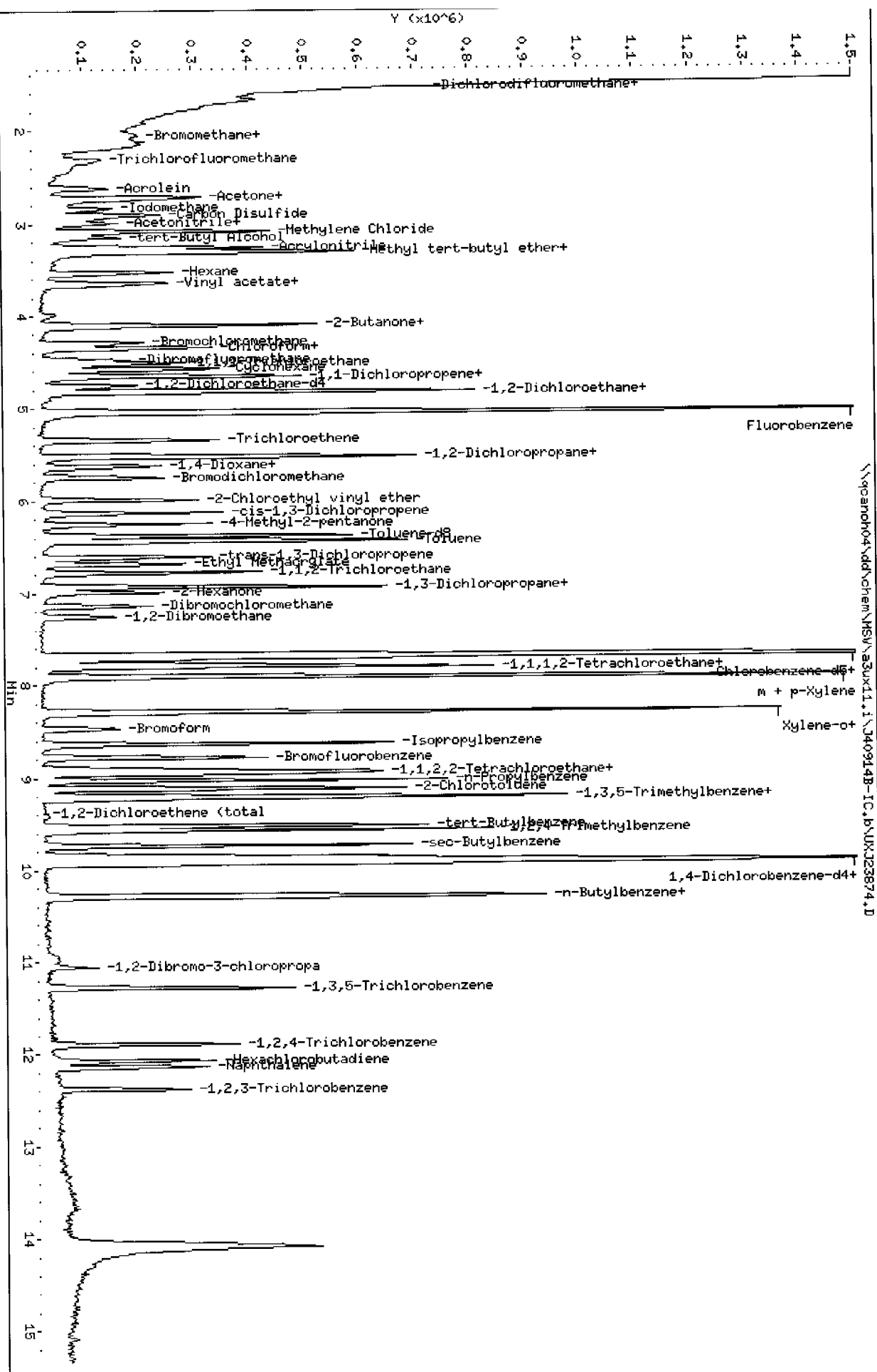
QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Instrument: aa30x11.i

Operator: 43592

Column diameter: 0.18
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STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23874.D
Lab Smp Id: 10NG-IC
Inj Date : 14-SEP-2004 15:19
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 10NG-IC
Misc Info : J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 1, 2
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:47 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 6 Calibration Sample, Level: 2
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ng)
*	1 Fluorobenzene	96	5.029	5.029 (1.000)	2293402	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1817387	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	990181	50.0000		
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.889)	106274	10.0000	9.953	
\$	5 1,2-Dichloroethane-d4	65	4.745	4.745 (0.944)	141757	10.0000	9.655	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	427652	10.0000	9.795	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	185658	10.0000	10.008	
8	Dichlorodifluoromethane	85	1.526	1.526 (0.304)	108953	10.0000	9.087	
9	Chloromethane	50	1.680	1.680 (0.334)	228871	10.0000	10.691	
10	Vinyl Chloride	62	1.763	1.763 (0.351)	146078	10.0000	9.976	
11	Bromomethane	94	2.047	2.047 (0.407)	75436	10.0000	10.861	
12	Chloroethane	64	2.118	2.118 (0.421)	113260	10.0000	10.524	
13	Trichlorofluoromethane	101	2.307	2.307 (0.459)	142867	10.0000	9.308	
15	Acrolein	56	2.615	2.615 (0.520)	140593	100.000	98.140	
16	Acetone	43	2.722	2.722 (0.541)	128042	20.0000	20.736	
17	1,1-Dichloroethene	96	2.710	2.710 (0.539)	105536	10.0000	10.290	
18	Freon-113	151	2.745	2.745 (0.546)	76261	10.0000	11.396	

Compounds	QUANT SIG							AMOUNTS	
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)	
		====	==	=====	=====	=====	=====	=====	=====
19 Iodomethane		142	2.840	2.840 (0.565)		159856	10.0000	10.545	
20 Carbon Disulfide		76	2.899	2.899 (0.577)		400515	10.0000	10.454	
21 Methylene Chloride		84	3.077	3.077 (0.612)		251128	10.0000	10.978	
22 Acetonitrile		41	2.946	2.946 (0.586)		138464	100.000	101.73	
23 Acrylonitrile		53	3.254	3.254 (0.647)		439101	100.000	103.74	
24 Methyl tert-butyl ether		73	3.301	3.301 (0.657)		308600	10.0000	9.385	
25 trans-1,2-Dichloroethene		96	3.313	3.313 (0.659)		126088	10.0000	10.363	
26 Hexane		86	3.538	3.538 (0.704)		22298	10.0000	11.260	
27 Vinyl acetate		43	3.656	3.656 (0.727)		175571	10.0000	9.018	
28 1,1-Dichloroethane		63	3.633	3.633 (0.722)		226350	10.0000	10.198	
29 tert-Butyl Alcohol		59	3.148	3.148 (0.626)		185158	200.000	207.57	
30 2-Butanone		43	4.082	4.082 (0.812)		117242	20.0000	18.664	
M 31 1,2-Dichloroethene (total)		96				261363	20.0000	20.945	
32 cis-1,2-dichloroethene		96	4.094	4.094 (0.814)		135275	10.0000	10.582	
33 2,2-Dichloropropane		77	4.106	4.106 (0.816)		141153	10.0000	10.553	
34 Bromochloromethane		128	4.283	4.283 (0.852)		65133	10.0000	11.003	
35 Chloroform		83	4.343	4.343 (0.864)		231979	10.0000	10.397	
36 Tetrahydrofuran		42	4.331	4.331 (0.861)		38288	10.0000	11.006	
37 1,1,1-Trichloroethane		97	4.508	4.508 (0.896)		171781	10.0000	10.194	
38 1,1-Dichloropropene		75	4.650	4.650 (0.925)		163880	10.0000	10.435	
39 Carbon Tetrachloride		117	4.650	4.650 (0.925)		134205	10.0000	10.143	
40 1,2-Dichloroethane		62	4.816	4.816 (0.958)		194488	10.0000	10.668	
41 Benzene		78	4.816	4.816 (0.958)		545878	10.0000	10.338	
42 Trichloroethene		130	5.337	5.337 (1.061)		121044	10.0000	10.045	
43 1,2-Dichloropropane		63	5.526	5.526 (1.099)		138162	10.0000	10.607	
44 1,4-Dioxane		88	5.621	5.621 (1.118)		53454	500.000	478.78 (A)	
45 Dibromomethane		93	5.621	5.621 (1.118)		76444	10.0000	10.591	
46 Bromodichloromethane		83	5.751	5.751 (1.144)		176533	10.0000	10.303	
47 2-Chloroethyl vinyl ether		63	5.987	5.987 (1.191)		135749	20.0000	18.535	
48 cis-1,3-Dichloropropene		75	6.129	6.129 (1.219)		218946	10.0000	10.444	
49 4-Methyl-2-pentanone		43	6.248	6.248 (1.242)		216822	20.0000	19.234	
50 Toluene		91	6.437	6.437 (0.838)		563016	10.0000	10.373	
51 trans-1,3-Dichloropropene		75	6.603	6.603 (0.860)		193073	10.0000	9.921	
52 Ethyl Methacrylate		69	6.674	6.674 (0.869)		163654	10.0000	9.774	
53 1,1,2-Trichloroethane		97	6.768	6.768 (0.881)		114065	10.0000	10.360	
54 1,3-Dichloropropane		76	6.922	6.922 (0.901)		215113	10.0000	10.830	
55 Tetrachloroethene		164	6.934	6.934 (0.903)		95253	10.0000	10.832	
56 2-Hexanone		43	6.981	6.981 (0.909)		173068	20.0000	19.692	
57 Dibromochloromethane		129	7.135	7.135 (0.929)		121301	10.0000	10.330	
58 1,2-Dibromoethane		107	7.254	7.254 (0.945)		114994	10.0000	10.601	
59 Chlorobenzene		112	7.703	7.703 (1.003)		377942	10.0000	10.754	
60 1,1,1,2-Tetrachloroethane		131	7.774	7.774 (1.012)		125087	10.0000	10.245	
61 Ethylbenzene		106	7.798	7.798 (1.015)		184529	10.0000	10.397	
62 m + p-Xylene		106	7.904	7.904 (1.029)		470158	20.0000	20.568	
M 63 Xylenes (total)		106				701783	30.0000	30.890	
64 Xylene-o		106	8.283	8.283 (1.079)		231625	10.0000	10.322	
65 Styrene		104	8.295	8.295 (1.080)		406292	10.0000	10.124	

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)
66 Bromoform	====	173	8.472	8.472 (1.103)		81750	10.0000	10.242
67 Isopropylbenzene	====	105	8.626	8.626 (1.123)		495185	10.0000	9.908
68 1,1,2,2-Tetrachloroethane	====	83	8.898	8.898 (0.898)		154970	10.0000	10.382
69 1,4-Dichloro-2-butene	====	53	8.957	8.957 (0.904)		47422	10.0000	9.656
70 1,2,3-Trichloropropane	====	110	8.946	8.946 (0.903)		49915	10.0000	10.191
71 Bromobenzene	====	156	8.934	8.934 (0.902)		154286	10.0000	10.444
72 n-Propylbenzene	====	120	9.028	9.028 (0.912)		140117	10.0000	10.173
73 2-Chlorotoluene	====	126	9.111	9.111 (0.920)		137109	10.0000	10.052
74 1,3,5-Trimethylbenzene	====	105	9.194	9.194 (0.928)		449094	10.0000	9.798
75 4-Chlorotoluene	====	126	9.218	9.218 (0.931)		150151	10.0000	10.304
76 tert-Butylbenzene	====	119	9.514	9.514 (0.961)		384632	10.0000	10.359
77 1,2,4-Trimethylbenzene	====	105	9.561	9.561 (0.965)		494544	10.0000	10.108
78 sec-Butylbenzene	====	105	9.727	9.727 (0.982)		506228	10.0000	10.072
79 4-Isopropyltoluene	====	119	9.869	9.869 (0.996)		417405	10.0000	9.982
80 1,3-Dichlorobenzene	====	146	9.845	9.845 (0.994)		280475	10.0000	10.284
81 1,4-Dichlorobenzene	====	146	9.928	9.928 (1.002)		298356	10.0000	10.456
82 n-Butylbenzene	====	91	10.271	10.271 (1.037)		380534	10.0000	10.187
83 1,2-Dichlorobenzene	====	146	10.295	10.295 (1.039)		278506	10.0000	10.368
84 1,2-Dibromo-3-chloropropane	====	157	11.064	11.064 (1.117)		24938	10.0000	9.866
85 1,2,4-Trichlorobenzene	====	180	11.892	11.892 (1.201)		120755	10.0000	10.559
86 Hexachlorobutadiene	====	225	12.070	12.070 (1.219)		62649	10.0000	9.935
87 Naphthalene	====	128	12.129	12.129 (1.225)		251285	10.0000	9.310
88 1,2,3-Trichlorobenzene	====	180	12.377	12.377 (1.250)		87516	10.0000	10.671
98 Cyclohexane	====	56	4.567	4.567 (0.908)		177719	10.0000	10.533
143 Methyl Acetate	====	43	2.994	2.994 (0.595)		199578	20.0000	21.155
144 Methylcyclohexane	====	83	5.514	5.514 (1.096)		144963	10.0000	10.239
141 1,3,5-Trichlorobenzene	====	180	11.277	11.277 (1.139)		157787	10.0000	10.452

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

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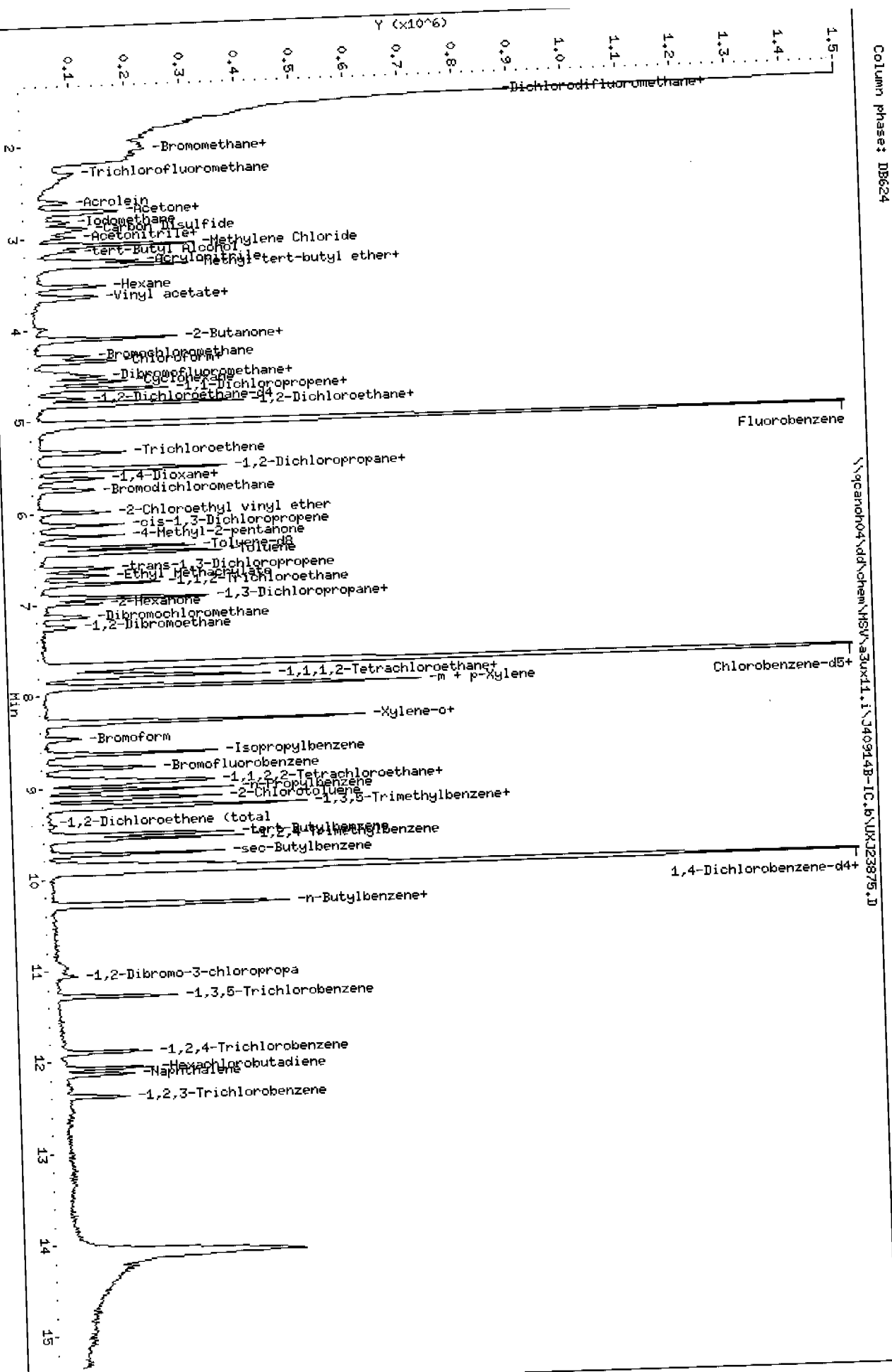
Instrument: a3ux11, i

Sample Info: 5NG-IC

Galilee 2008: 5.0

כטבָּה וְזִקְנָה יְהוָה

ଲୋକାନ୍ତିର ପାଦମଣି



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23875.D
Lab Smp Id: 5NG-IC
Inj Date : 14-SEP-2004 15:41
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 5NG-IC
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,1,1
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 12:47 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 7 Calibration Sample, Level: 1
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ng)
*	1 Fluorobenzene	96	5.029	5.029 (1.000)	2251983	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1826599	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	937868	50.0000		
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.889)	51569	5.00000	4.918	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.946)	70414	5.00000	4.884	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	210171	5.00000	4.789	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	88368	5.00000	4.739	
8	Dichlorodifluoromethane	85	1.526	1.526 (0.304)	64938	5.00000	5.516	
9	Chloromethane	50	1.668	1.668 (0.332)	125378	5.00000	5.964	
10	Vinyl Chloride	62	1.763	1.763 (0.351)	72850	5.00000	5.066	
11	Bromomethane	94	2.035	2.035 (0.405)	38866	5.00000	5.699	
12	Chloroethane	64	2.118	2.118 (0.421)	52345	5.00000	4.953	
13	Trichlorofluoromethane	101	2.307	2.307 (0.459)	86493	5.00000	5.739	
15	Acrolein	56	2.615	2.615 (0.520)	72041	50.0000	51.213	
16	Acetone	43	2.722	2.722 (0.541)	72580	10.0000	9.714	
17	1,1-Dichloroethene	96	2.710	2.710 (0.539)	55428	5.00000	5.504	
18	Freon-113	151	2.745	2.745 (0.546)	37991	5.00000	5.276	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23875.D
 Report Date: 15-Sep-2004 12:48

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)	ON-COL (ng)
19 Iodomethane		142	2.828	2.828 (0.562)		70148	5.00000	4.712
20 Carbon Disulfide		76	2.899	2.899 (0.577)		205480	5.00000	5.462
21 Methylene Chloride		84	3.077	3.077 (0.612)		169409	5.00000	4.472
22 Acetonitrile		41	2.946	2.946 (0.586)		74099	50.0000	55.440
23 Acrylonitrile		53	3.254	3.254 (0.647)		205768	50.0000	49.508
24 Methyl tert-butyl ether		73	3.301	3.301 (0.657)		147953	5.00000	4.582
25 trans-1,2-Dichloroethene		96	3.313	3.313 (0.659)		63645	5.00000	5.327
26 Hexane		86	3.526	3.526 (0.701)		11663	5.00000	5.789
27 Vinyl acetate		43	3.656	3.656 (0.727)		93501	5.00000	4.891
28 1,1-Dichloroethane		63	3.633	3.633 (0.722)		115670	5.00000	5.307
29 tert-Butyl Alcohol		59	3.148	3.148 (0.626)		92424	100.000	105.52
30 2-Butanone		43	4.094	4.094 (0.814)		71588	10.0000	11.606
M 31 1,2-Dichloroethene (total)		96				128452	10.0000	10.490
32 cis-1,2-dichloroethene		96	4.106	4.106 (0.816)		64807	5.00000	5.163
33 2,2-Dichloropropane		77	4.106	4.106 (0.816)		66643	5.00000	5.074
34 Bromochloromethane		128	4.295	4.295 (0.854)		28755	5.00000	4.947
35 Chloroform		83	4.343	4.343 (0.864)		114878	5.00000	5.243
36 Tetrahydrofuran		42	4.331	4.331 (0.861)		22560	5.00000	6.678
37 1,1,1-Trichloroethane		97	4.508	4.508 (0.896)		88473	5.00000	5.347
38 1,1-Dichloropropene		75	4.650	4.650 (0.925)		78641	5.00000	5.099
39 Carbon Tetrachloride		117	4.650	4.650 (0.925)		68204	5.00000	5.250
40 1,2-Dichloroethane		62	4.816	4.816 (0.958)		90135	5.00000	5.035
41 Benzene		78	4.816	4.816 (0.958)		283739	5.00000	5.472
42 Trichloroethene		130	5.349	5.349 (1.064)		61441	5.00000	5.192
43 1,2-Dichloropropane		63	5.514	5.514 (1.096)		66815	5.00000	5.224
44 1,4-Dioxane		88	5.621	5.621 (1.118)		27287	250.000	248.90 (A)
45 Dibromomethane		93	5.621	5.621 (1.118)		34422	5.00000	4.857
46 Bromodichloromethane		83	5.751	5.751 (1.144)		86009	5.00000	5.112
47 2-Chloroethyl vinyl ether		63	5.988	5.988 (1.191)		62831	10.0000	8.737
48 cis-1,3-Dichloropropene		75	6.129	6.129 (1.219)		97774	5.00000	4.750
49 4-Methyl-2-pentanone		43	6.248	6.248 (1.242)		106568	10.0000	9.627
50 Toluene		91	6.437	6.437 (0.838)		271647	5.00000	4.980
51 trans-1,3-Dichloropropene		75	6.603	6.603 (0.860)		92511	5.00000	4.730
52 Ethyl Methacrylate		69	6.686	6.686 (0.871)		71117	5.00000	4.226
53 1,1,2-Trichloroethane		97	6.780	6.780 (0.883)		55141	5.00000	4.983
54 1,3-Dichloropropane		76	6.922	6.922 (0.901)		104493	5.00000	5.012
55 Tetrachloroethene		164	6.934	6.934 (0.903)		46594	5.00000	5.272
56 2-Hexanone		43	6.993	6.993 (0.911)		81226	10.0000	9.196
57 Dibromochloromethane		129	7.135	7.135 (0.929)		56564	5.00000	4.793
58 1,2-Dibromoethane		107	7.242	7.242 (0.943)		49174	5.00000	4.510
59 Chlorobenzene		112	7.703	7.703 (1.003)		176312	5.00000	4.992
60 1,1,1,2-Tetrachloroethane		131	7.774	7.774 (1.012)		61700	5.00000	5.028
61 Ethylbenzene		106	7.798	7.798 (1.015)		84068	5.00000	4.713
62 m + p-Xylene		106	7.904	7.904 (1.029)		221783	10.0000	9.654
M 63 Xylenes (total)		106				325206	15.0000	14.239
64 Xylene-o		106	8.283	8.283 (1.079)		103423	5.00000	4.586
65 Styrene		104	8.295	8.295 (1.080)		183225	5.00000	4.543

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ238\5.D
 Report Date: 15-Sep-2004 12:48

Compounds	QUANT SIG	MASS	AMOUNTS				
			RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
		====	==	=====	=====	=====	=====
66 Bromoform		173	8.472	8.472 (1.103)		36774	5.00000 4.584
67 Isopropylbenzene		105	8.626	8.626 (1.123)		245674	5.00000 4.891
68 1,1,2,2-Tetrachloroethane		83	8.898	8.898 (0.898)		69472	5.00000 4.914
69 1,4-Dichloro-2-butene		53	8.958	8.958 (0.904)		21722	5.00000 4.670
70 1,2,3-Trichloropropane		110	8.946	8.946 (0.903)		23727	5.00000 5.114
71 Bromobenzene		156	8.934	8.934 (0.902)		69868	5.00000 4.993
72 n-Propylbenzene		120	9.029	9.029 (0.912)		67269	5.00000 5.156
73 2-Chlorotoluene		126	9.111	9.111 (0.920)		67172	5.00000 5.199
74 1,3,5-Trimethylbenzene		105	9.194	9.194 (0.928)		211928	5.00000 4.882
75 4-Chlorotoluene		126	9.218	9.218 (0.931)		71891	5.00000 5.209
76 tert-Butylbenzene		119	9.514	9.514 (0.961)		177303	5.00000 5.041
77 1,2,4-Trimethylbenzene		105	9.561	9.561 (0.965)		218676	5.00000 4.719
78 sec-Butylbenzene		105	9.727	9.727 (0.982)		249496	5.00000 5.241
79 4-Isopropyltoluene		119	9.869	9.869 (0.996)		190215	5.00000 4.803
80 1,3-Dichlorobenzene		146	9.845	9.845 (0.994)		141050	5.00000 5.460
81 1,4-Dichlorobenzene		146	9.928	9.928 (1.002)		139945	5.00000 5.178
82 n-Butylbenzene		91	10.271	10.271 (1.037)		174632	5.00000 5.343
83 1,2-Dichlorobenzene		146	10.295	10.295 (1.039)		135942	5.00000 4.952
84 1,2-Dibromo-3-chloropropane		157	11.064	11.064 (1.117)		11856	5.00000 4.882
85 1,2,4-Trichlorobenzene		180	11.892	11.892 (1.201)		52887	5.00000 4.872
86 Hexachlorobutadiene		225	12.070	12.070 (1.219)		36648	5.00000 4.386
87 Naphthalene		128	12.129	12.129 (1.225)		112143	5.00000 4.626
88 1,2,3-Trichlorobenzene		180	12.377	12.377 (1.250)		35937	5.00000 5.095
98 Cyclohexane		56	4.568	4.568 (0.908)		84417	5.00000 10.849
143 Methyl Acetate		43	2.994	2.994 (0.595)		100499	10.0000 5.612
144 Methylcyclohexane		83	5.514	5.514 (1.096)		78015	5.00000 5.624
141 1,3,5-Trichlorobenzene		180	11.277	11.277 (1.139)		80418	5.00000

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

STL North Canton

RECOVERY REPORT

Client Name:
 Sample Matrix: LIQUID
 Lab Smp Id: ICV
 Level: LOW
 Data Type: MS DATA
 SpikeList File: plexus-ck.spk
 Sublist File: 2-8260.SUB
 Method File: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
 Misc Info: J40914B-IC, 8260LLUX11, 2-8260.SUB, 43582, 3

Client SDG: SDGa00594
 Fraction: VOA
 Operator: 43582
 SampleType: METHSPIKE
 Quant Type: ISTD

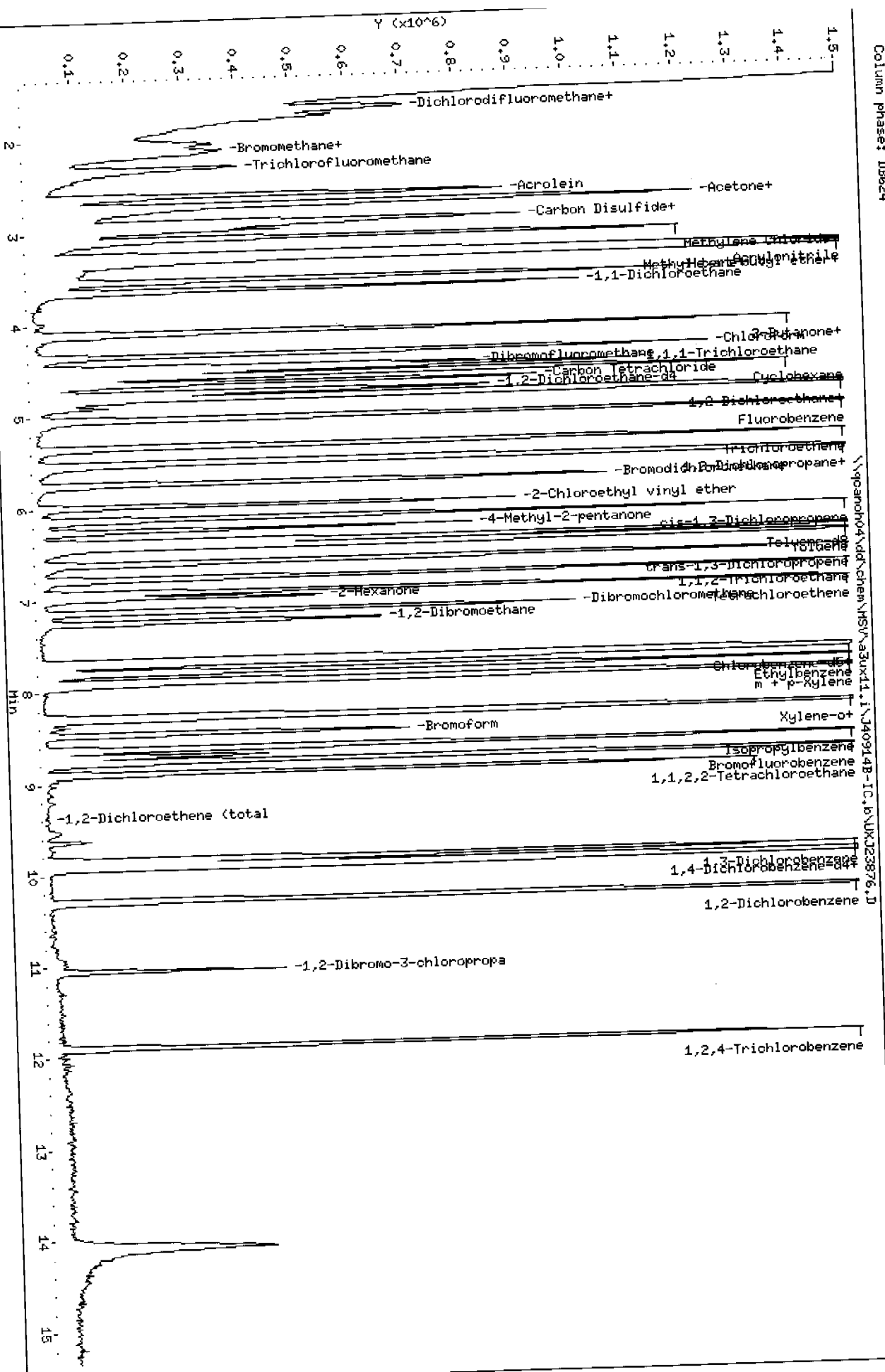
SPIKE COMPOUND	CONC ADDED ug/L	CONC RECOVERED ug/L	% RECOVERED	LIMITS
17 1,1-Dichloroethene	10.000	9.507	95.07	45-155
42 Trichloroethene	10.000	9.342	93.42	45-155
59 Chlorobenzene	10.000	9.443	94.43	45-155
50 Toluene	10.000	9.460	94.60	45-155
41 Benzene	10.000	9.294	92.94	45-155
16 Acetone	10.000	6.134	61.34	45-155
20 Carbon Disulfide	10.000	9.433	94.33	45-155
9 Chloromethane	10.000	8.196	81.96	45-155
11 Bromomethane	10.000	8.864	88.64	45-155
10 Vinyl Chloride	10.000	8.534	85.34	45-155
12 Chloroethane	10.000	8.736	87.36	45-155
21 Methylene Chloride	10.000	9.642	96.42	45-155
28 1,1-Dichloroethane	10.000	9.526	95.26	45-155
M	31 1,2-Dichloroethene	20.000	18.957	94.78
	35 Chloroform	10.000	9.419	94.19
	40 1,2-Dichloroethane	10.000	9.839	98.39
	30 2-Butanone	10.000	8.378	83.78
	37 1,1,1-Trichloroeth	10.000	9.100	91.00
	39 Carbon Tetrachlori	10.000	9.159	91.59
	46 Bromodichlorometha	10.000	9.513	95.13
	43 1,2-Dichloropropan	10.000	9.463	94.63
	48 cis-1,3-Dichloropr	10.000	9.690	96.90
	57 Dibromochlorometha	10.000	10.024	100.24
	53 1,1,2-Trichloroeth	10.000	9.656	96.56
	51 trans-1,3-Dichloro	10.000	9.392	93.92
	66 Bromoform	10.000	9.919	99.19
	49 4-Methyl-2-pentano	10.000	9.709	97.09
	56 2-Hexanone	10.000	8.736	87.36
	55 Tetrachloroethene	10.000	9.092	90.92
	68 1,1,2,2-Tetrachlor	10.000	10.303	103.03
	61 Ethylbenzene	10.000	9.427	94.27
	65 Styrene	10.000	9.617	96.17
	M 63 Xylenes (total)	30.000	28.176	93.92
				45-155

SPIKE COMPOUND	CONC ADDED ug/L	CONC RECOVERED ug/L	% RECOVERED	LIMITS
32 cis-1,2-dichloroet	10.000	9.524	95.24	45-155
25 trans-1,2-Dichloro	10.000	9.432	94.32	45-155
8 Dichlorodifluorome	10.000	9.117	91.17	45-155
13 Trichlorofluoromet	10.000	9.634	96.34	45-155
18 Freon-113	10.000	11.317	113.17	45-155
24 Methyl tert-butyl	10.000	9.278	92.78	45-155
58 1,2-Dibromoethane	10.000	9.902	99.02	45-155
67 Isopropylbenzene	10.000	9.750	97.50	45-155
80 1,3-Dichlorobenzen	10.000	9.577	95.77	45-155
81 1,4-Dichlorobenzen	10.000	9.879	98.79	45-155
83 1,2-Dichlorobenzen	10.000	9.570	95.70	45-155
84 1,2-Dibromo-3-chlo	10.000	10.335	103.35	45-155
85 1,2,4-Trichloroben	10.000	9.534	95.34	45-155
98 Cyclohexane	10.000	9.159	91.59	45-155
143 Methyl Acetate	10.000	9.302	93.02	45-155
144 Methylcyclohexane	10.000	8.960	89.60	45-155

SURROGATE COMPOUND	CONC ADDED ug/L	CONC RECOVERED ug/L	% RECOVERED	LIMITS
\$ 4 Dibromofluorometha	10.000	9.507	95.07	73-122
\$ 5 1,2-Dichloroethane	10.000	9.840	98.40	61-128
\$ 6 Toluene-d8	10.000	9.847	98.47	76-110
\$ 7 Bromofluorobenzene	10.000	10.152	101.52	74-116

Instrument: 3uX11.i

Operator: 43582
 Column diameter: 0.18
 \\pcanonh04\dd\chem\MSV\3uX11.i\\J40914B-1C.b\\UKJ23876.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23876.D
Report Date: 15-Sep-2004 13:02

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJ23876.D
Lab Smp Id: ICV
Inj Date : 14-SEP-2004 16:04
Operator : 43582 Inst ID: a3ux11.i
Smp Info : ICV
Misc Info : J40914B-IC,8260LLUX11,2-8260.SUB,43582,3
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\8260LLUX11.m
Meth Date : 15-Sep-2004 13:00 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 8 QC Sample: METHSPIKE
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ug/L)
*	1 Fluorobenzene	96	5.029	5.029 (1.000)	2348288	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1882362	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	979729	50.0000		
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.889)	519715	47.5349	9.507	
\$	5 1,2-Dichloroethane-d4	65	4.745	4.757 (0.944)	739678	49.1997	9.840	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	2226509	49.2353	9.847	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	975336	50.7609	10.152	
8	Dichlorodifluoromethane	85	1.527	1.526 (0.304)	559620	45.5836	9.117	
9	Chloromethane	50	1.680	1.668 (0.334)	898254	40.9791	8.196	
10	Vinyl Chloride	62	1.775	1.763 (0.353)	639778	42.6691	8.534	
11	Bromomethane	94	2.047	2.035 (0.407)	315198	44.3200	8.864	
12	Chloroethane	64	2.118	2.118 (0.421)	481308	43.6780	8.736	
13	Trichlorofluoromethane	101	2.308	2.307 (0.459)	756987	48.1676	9.634	
15	Acrolein	56	2.615	2.615 (0.520)	1028164	700.930	140.18	
16	Acetone	43	2.722	2.722 (0.541)	180849	30.6678	6.134	
17	1,1-Dichloroethene	96	2.710	2.710 (0.539)	499203	47.5355	9.507	
18	Freon-113	151	2.734	2.745 (0.544)	373218	56.5832	11.317	
19	Iodomethane	142		Compound Not Detected.				

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40914B-IC.b\UXJZ5810.D
 Report Date: 15-Sep-2004 13:02

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN	FINAL
							(ng)	(ug/L)
20 Carbon Disulfide		76	2.899	2.899 (0.577)	1.850176	47.1646	9.433	
21 Methylene Chloride		84	3.077	3.077 (0.612)	719449	48.2118	9.642	
22 Acetonitrile		41	2.935	2.946 (0.584)	665528	477.523	95.505	
23 Acrylonitrile		53	3.254	3.254 (0.647)	2230444	514.640	102.93	
24 Methyl tert-butyl ether		73	3.302	3.301 (0.657)	1561889	46.3911	9.278	
25 trans-1,2-Dichloroethene		96	3.313	3.313 (0.659)	587539	47.1615	9.432	
26 Hexane		86	3.526	3.526 (0.701)	107938	53.4083	10.682	
27 Vinyl acetate		43	3.526	3.656 (0.701)	387039	19.4150	3.883	
28 1,1-Dichloroethane		63	3.633	3.633 (0.722)	1082468	47.6309	9.526	
29 tert-Butyl Alcohol		59	2.994	3.148 (0.595)	39015	42.7151	8.543	
30 2-Butanone		43	4.083	4.094 (0.812)	269449	41.8924	8.378	
M 31 1,2-Dichloroethene (total)		96			1210887	94.7835	18.957	
32 cis-1,2-dichloroethene		96	4.094	4.106 (0.814)	623348	47.6220	9.524	
33 2,2-Dichloropropane		77		Compound Not Detected.				
34 Bromochloromethane		128		Compound Not Detected.				
35 Chloroform		83	4.343	4.343 (0.864)	1075879	47.0934	9.419	
36 Tetrahydrofuran		42	4.083	4.331 (0.812)	19114	5.46454	1.093	
37 1,1,1-Trichloroethane		97	4.509	4.508 (0.896)	785098	45.5027	9.100	
38 1,1-Dichloropropene		75		Compound Not Detected.				
39 Carbon Tetrachloride		117	4.662	4.650 (0.927)	620420	45.7962	9.159	
40 1,2-Dichloroethane		62	4.816	4.816 (0.958)	918291	49.1951	9.839	
41 Benzene		78	4.816	4.816 (0.958)	2512537	46.4726	9.294	
42 Trichloroethene		130	5.337	5.349 (1.061)	576326	46.7078	9.342	
43 1,2-Dichloropropane		63	5.526	5.514 (1.099)	631042	47.3158	9.463	
44 1,4-Dioxane		88		Compound Not Detected.				
45 Dibromomethane		93		Compound Not Detected.				
46 Bromodichloromethane		83	5.751	5.751 (1.144)	834464	47.5633	9.513	
47 2-Chloroethyl vinyl ether		63	5.988	5.988 (1.191)	393567	52.4826	10.496	
48 cis-1,3-Dichloropropene		75	6.130	6.129 (1.219)	1040015	48.4491	9.690	
49 4-Methyl-2-pentanone		43	6.248	6.248 (1.242)	560341	48.5456	9.709	
50 Toluene		91	6.437	6.437 (0.838)	2659243	47.3027	9.460	
51 trans-1,3-Dichloropropene		75	6.603	6.603 (0.860)	946487	46.9582	9.392	
52 Ethyl Methacrylate		69		Compound Not Detected.				
53 1,1,2-Trichloroethane		97	6.769	6.780 (0.881)	550573	48.2825	9.656	
54 1,3-Dichloropropane		76		Compound Not Detected.				
55 Tetrachloroethene		164	6.934	6.934 (0.903)	414030	45.4582	9.092	
56 2-Hexanone		43	6.982	6.993 (0.909)	397634	43.6825	8.736	
57 Dibromochloromethane		129	7.135	7.135 (0.929)	609608	50.1213	10.024	
58 1,2-Dibromoethane		107	7.254	7.242 (0.945)	556274	49.5113	9.902	
59 Chlorobenzene		112	7.703	7.703 (1.003)	1718695	47.2166	9.443	
60 1,1,1,2-Tetrachloroethane		131		Compound Not Detected.				
61 Ethylbenzene		106	7.798	7.798 (1.015)	866451	47.1356	9.427	
62 m + p-Xylene		106	7.905	7.904 (1.029)	2209377	93.3186	18.664	
M 63 Xylenes (total)		106			3314793	140.878	28.176	
64 Xylene-o		104	8.283	8.283 (1.079)	1105416	47.5597	9.512	
65 Styrene		104	8.295	8.295 (1.080)	1998604	48.0849	9.617	
66 Bromoform		173	8.473	8.472 (1.103)	410026	49.5965	9.919	

Compounds	QUANT SIG	MASS	CONCENTRATIONS				
			RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
67 Isopropylbenzene		105	8.626	8.626 (1.123)	2523492	48.7480	9.750
68 1,1,2,2-Tetrachloroethane		83	8.899	8.898 (0.898)	760843	51.5174	10.303
69 1,4-Dichloro-2-butene		53	Compound Not Detected.				
70 1,2,3-Trichloropropane		110	Compound Not Detected.				
71 Bromobenzene		156	Compound Not Detected.				
72 n-Propylbenzene		120	Compound Not Detected.				
73 2-Chlorotoluene		126	Compound Not Detected.				
74 1,3,5-Trimethylbenzene		105	Compound Not Detected.				
75 4-Chlorotoluene		126	Compound Not Detected.				
76 tert-Butylbenzene		119	Compound Not Detected.				
77 1,2,4-Trimethylbenzene		105	Compound Not Detected.				
78 sec-Butylbenzene		105	Compound Not Detected.				
79 4-Isopropyltoluene		119	Compound Not Detected.				
80 1,3-Dichlorobenzene		146	9.845	9.845 (0.994)	1292290	47.8872	9.577
81 1,4-Dichlorobenzene		146	9.928	9.928 (1.002)	1394610	49.3955	9.879
82 n-Butylbenzene		91	Compound Not Detected.				
83 1,2-Dichlorobenzene		146	10.295	10.295 (1.039)	1271863	47.8511	9.570
84 1,2-Dibromo-3-chloropropane		157	11.052	11.064 (1.116)	129232	51.6742	10.335
85 1,2,4-Trichlorobenzene		180	11.892	11.892 (1.201)	539429	47.6703	9.534
86 Hexachlorobutadiene		225	Compound Not Detected.				
87 Naphthalene		128	Compound Not Detected.				
88 1,2,3-Trichlorobenzene		180	Compound Not Detected.				
98 Cyclohexane		56	4.568	4.568 (0.908)	791178	45.7968	9.159
143 Methyl Acetate		43	2.994	2.994 (0.595)	449301	46.5123	9.302
144 Methylcyclohexane		83	5.514	5.514 (1.096)	649471	44.7998	8.960
141 1,3,5-Trichlorobenzene		180	Compound Not Detected.				

Calibration History

Method : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Start Cal Date: 16-AUG-2004 16:18
End Cal Date : 14-SEP-2004 15:41
Last Cal Level: 1
Last Cal Type : Initial Calibration

Initial Calibration

Injection Date	Sublist	Calibration File
Cal Level: 1 , Cal Amount: 5.000		
14-SEP-2004 15:41	2-8260	UXJ23875.D
16-AUG-2004 18:11	3-IX	UXJ23214.D
Cal Level: 2 , Cal Amount: 10.000		
14-SEP-2004 15:19	2-8260	UXJ23874.D
16-AUG-2004 17:48	3-IX	UXJ23213.D
Cal Level: 3 , Cal Amount: 25.000		
14-SEP-2004 14:57	2-8260	UXJ23873.D
16-AUG-2004 17:26	3-IX	UXJ23212.D
Cal Level: 4 , Cal Amount: 50.000		
14-SEP-2004 14:33	2-8260	UXJ23872.D
16-AUG-2004 17:03	3-IX	UXJ23211.D
Cal Level: 5 , Cal Amount: 100.00		
14-SEP-2004 14:10	2-8260	UXJ23871.D
16-AUG-2004 16:40	3-IX	UXJ23210.D
Cal Level: 6 , Cal Amount: 200.00		
14-SEP-2004 13:48	2-8260	UXJ23870.D
16-AUG-2004 16:18	3-IX	UXJ23209.D

Continuing Calibration

29-SEP-2004 07:50	2-8260	UXJ24189.D
29-SEP-2004 08:14	3-IX	UXJ24190.D

Report Date: 09/30/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24189.D
Analysis Type: WATER

Injection Date: 29-SEP-2004 07:50
Lab Sample ID: 50NG-CC
Method File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\J40929A

COMPOUND	EXPECTED	MEASURED	%D	MAX
	CONC.	CONC.		
0 Chlorobenzene	50.0000	51.7337	3.5	50.0
0 Bromodichloromethane	50.0000	49.2014	1.6	50.0
0 1,1,2,2-Tetrachloroethane	50.0000	52.4691	4.9	50.0
0 Bromoform	50.0000	52.3396	4.7	50.0
0 Styrene	50.0000	56.1883	12.4	50.0
0 Xylene-o	50.0000	54.4505	8.9	50.0
0 Xylenes (total)	150.0000	162.3570	8.2	50.0
0 2-Hexanone	100.0000	98.4586	1.5	50.0
0 Chloromethane	50.0000	56.2159	12.4	50.0
0 Vinyl Chloride	50.0000	52.8242	5.6	20.0
0 Bromomethane	50.0000	39.2952	21.4	50.0
0 Chloroethane	50.0000	57.3743	14.7	50.0
0 1,1-Dichloroethane	50.0000	49.6485	0.7	50.0
0 Tetrachloroethene	50.0000	51.1363	2.3	50.0
0 Acetone	100.0000	84.8478	15.2	50.0
0 1,1-Dichloroethene	50.0000	50.4067	0.8	20.0
0 m + p-Xylene	100.0000	107.9065	7.9	50.0
0 Ethylbenzene	50.0000	51.3446	2.7	20.0
0 Carbon Disulfide	50.0000	52.8300	5.7	50.0
0 Methylene Chloride	50.0000	64.7808	29.6	50.0
0 1,2-Dichloropropane	50.0000	49.2037	1.6	20.0
0 1,1,2-Trichloroethane	50.0000	53.1927	6.4	50.0
0 Dibromochloromethane	50.0000	50.7705	1.5	50.0
0 trans-1,2-Dichloroethene	50.0000	49.8167	0.4	50.0
0 trans-1,3-Dichloropropene	50.0000	43.6062	12.8	50.0
0 cis-1,3-Dichloropropene	50.0000	45.8697	8.3	50.0
0 Chloroform	50.0000	48.9534	2.1	20.0
0 Toluene	50.0000	49.4329	1.1	20.0
0 2-Butanone	100.0000	82.7906	17.2	50.0
0 1,2-Dichloroethene (total)	100.0000	96.7504	3.2	50.0
0 cis-1,2-dichloroethene	50.0000	46.9337	6.1	50.0
0 4-Methyl-2-pentanone	100.0000	113.5068	13.5	50.0
0 1,2-Dichloroethane	50.0000	49.5922	0.8	50.0
0 Trichloroethene	50.0000	46.8037	6.4	50.0
0 1,1,1-Trichloroethane	50.0000	43.9125	12.2	50.0
0 Carbon Tetrachloride	50.0000	49.0579	1.9	50.0
0 Benzene	50.0000	50.4168	0.8	50.0
38 Dichlorodifluoromethane	50.0000	68.4174	36.8	50.0
39 Trichlorofluoromethane	50.0000	42.3684	15.3	50.0

Report Date: 09/30/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24189.D
Analysis Type: WATER

Injection Date: 29-SEP-2004 07:50
Lab Sample ID: 50NG-CC
Method File: \\QCANOH04\\dd\\chem\\MSV\\a3ux11.i\\

COMPOUND	EXPECTED	MEASURED	%D	MAX
	CONC.	CONC.		
39 Chlorobenzene-d5	50.0000	50.0000	0.0	50.0
40 Acrolein	500.0000	584.1857	16.8	50.0
41 Acrylonitrile	500.0000	542.2128	8.4	50.0
42 Vinyl acetate	50.0000	45.7275	8.5	50.0
43 2-Chloroethyl vinyl ether	100.0000	95.0217	5.0	50.0
47 Freon-113	50.0000	61.1381	22.3	50.0
48 1,3-Dichlorobenzene	50.0000	49.5587	0.9	50.0
49 1,4-Dichlorobenzene	50.0000	49.3159	1.4	50.0
50 1,2-Dichlorobenzene	50.0000	48.8545	2.3	50.0
51 Acetonitrile	500.0000	635.9772	27.2	50.0
52 Iodomethane	50.0000	44.8218	10.4	50.0
59 1,4-Dioxane	2500.0000	2907.7908	16.3	50.0
60 Dibromomethane	50.0000	52.2437	4.5	50.0
62 Ethyl Methacrylate	50.0000	50.2399	0.5	50.0
63 1,2-Dibromoethane	50.0000	49.9206	0.2	50.0
64 1,1,1,2-Tetrachloroethane	50.0000	49.7466	0.5	50.0
65 1,2,3-Trichloropropane	50.0000	51.0593	2.1	50.0
66 1,4-Dichloro-2-butene	50.0000	47.7869	4.4	50.0
69 1,2-Dibromo-3-chloropropane	50.0000	36.2438	27.5	50.0
82 Methyl tert-butyl ether	50.0000	43.0090	14.0	50.0
84 Tetrahydrofuran	50.0000	45.6028	8.8	50.0
98 2,2-Dichloropropane	50.0000	37.2350	25.5	50.0
99 1,1-Dichloropropene	50.0000	49.9337	0.1	50.0
100 1,3-Dichloropropane	50.0000	49.4257	1.1	50.0
102 Bromobenzene	50.0000	51.7885	3.6	50.0
103 2-Chlorotoluene	50.0000	50.9425	1.9	50.0
104 n-Propylbenzene	50.0000	53.8098	7.6	50.0
105 4-Chlorotoluene	50.0000	53.0453	6.1	50.0
106 1,3,5-Trimethylbenzene	50.0000	51.4587	2.9	50.0
107 tert-Butylbenzene	50.0000	47.4989	5.0	50.0
108 1,2,4-Trimethylbenzene	50.0000	50.5932	1.2	50.0
109 sec-Butylbenzene	50.0000	48.3953	3.2	50.0
110 4-Isopropyltoluene	50.0000	48.7147	2.6	50.0
111 n-Butylbenzene	50.0000	45.4323	9.1	50.0
112 1,2,4-Trichlorobenzene	50.0000	25.3974	49.2	50.0
113 Naphthalene	50.0000	16.1837	67.6	50.0 <-
114 Hexachlorobutadiene	50.0000	41.1506	17.7	50.0
115 1,2,3-Trichlorobenzene	50.0000	18.9283	62.1	50.0 <-
124 tert-Butyl Alcohol	1000.0000	882.6330	11.7	50.0

Report Date: 09/30/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24189.D
Analysis Type: WATER

Injection Date: 29-SEP-2004 07:50
Lab Sample ID: 50NG-CC
Method File: \\QCANOHO4\\dd\\chem\\MSV\\a3ux11.i\\

COMPOUND	EXPECTED	MEASURED	%D	MAX
	CONC.	CONC.		
125 Hexane	50.0000	52.3522	4.7	20.0
127 Cyclohexane	50.0000	50.7918	1.6	50.0
128 Isopropylbenzene	50.0000	51.2812	2.6	50.0
130 Fluorobenzene	50.0000	50.0000	0.0	50.0
132 1,4-Dichlorobenzene-d4	50.0000	50.0000	0.0	50.0
133 Bromochloromethane	50.0000	51.7935	3.6	50.0
141 1,3,5-Trichlorobenzene	50.0000	39.6222	20.8	50.0
143 Methyl Acetate	100.0000	95.1989	4.8	50.0
144 Methylcyclohexane	50.0000	49.0461	1.9	50.0
22 Toluene-d8	50.0000	50.5004	1.0	50.0
32 Bromofluorobenzene	50.0000	53.3691	6.7	50.0
47 1,2-Dichloroethane-d4	50.0000	53.0223	6.0	50.0
131 Dibromofluoromethane	50.0000	52.4657	4.9	50.0

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24189.D
Report Date: 29-Sep-2004 08:40

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 29-SEP-2004 07:50
Lab File ID: UXJ24189.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m

COMPOUND	RRF	RF50	MIN	MAX
\$ 4 Dibromofluoromethane	0.23279	0.24427	0.010	4.9 50.0
\$ 5 1,2-Dichloroethane-d4	0.32011	0.33946	0.010	6.0 50.0
\$ 6 Toluene-d8	1.20120	1.21322	0.010	1.0 50.0
\$ 7 Bromofluorobenzene	0.51038	0.54477	0.010	6.7 50.0
8 Dichlorodifluoromethane	0.26140	0.35769	0.010	36.8 50.0
9 Chloromethane	0.46672	0.52474	0.100	12.4 50.0
10 Vinyl Chloride	0.31925	0.33729	0.010	5.6 20.0
11 Bromomethane	0.15143	0.11901	0.010	-21.4 50.0
12 Chloroethane	0.23463	0.26923	0.010	14.7 50.0
13 Trichlorofluoromethane	0.33462	0.28355	0.010	-15.3 50.0
15 Acrolein	0.03123	0.03649	0.010	16.8 50.0
16 Acetone	100	84.84780	0.010	15.2 50.0
17 1,1-Dichloroethene	0.22360	0.22542	0.010	0.8 20.0
18 Freon-113	50.00000	61.13812	0.010	-22.3 50.0
19 Iodomethane	0.33050	0.29627	0.010	-10.4 50.0
20 Carbon Disulfide	0.83525	0.88252	0.010	5.7 50.0
21 Methylene Chloride	50.00000	64.78084	0.010	-29.6 50.0
22 Acetonitrile	0.02967	0.03775	0.010	27.2 50.0
23 Acrylonitrile	0.09228	0.10007	0.010	8.4 50.0
24 Methyl tert-butyl ether	0.71686	0.61663	0.010	-14.0 50.0
25 trans-1,2-Dichloroethene	0.26526	0.26429	0.010	-0.4 50.0
26 Hexane	50.00000	52.35224	0.010	-4.7 20.0
27 Vinyl acetate	0.42446	0.38819	0.010	-8.5 50.0
28 1,1-Dichloroethane	0.48389	0.48049	0.100	-0.7 50.0
29 tert-Butyl Alcohol	0.01945	0.01717	0.010	-11.7 50.0
30 2-Butanone	0.13695	0.11338	0.010	-17.2 50.0
M 31 1,2-Dichloroethene (total)	0.27198	0.26295	0.010	-3.3 50.0
32 cis-1,2-dichloroethene	0.27870	0.26161	0.010	-6.1 50.0
33 2,2-Dichloropropane	0.29161	0.21716	0.010	-25.5 50.0
34 Bromochloromethane	0.12906	0.13369	0.010	3.6 50.0
35 Chloroform	0.48643	0.47625	0.010	-2.1 20.0
36 Tetrahydrofuran	50.00000	45.60279	0.010	8.8 50.0
37 1,1,1-Trichloroethane	0.36737	0.32264	0.010	-12.2 50.0
38 1,1-Dichloropropene	0.34240	0.34194	0.010	-0.1 50.0
39 Carbon Tetrachloride	0.28845	0.28302	0.010	-1.9 50.0
40 1,2-Dichloroethane	0.39745	0.39420	0.010	-0.8 50.0

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24189.D
Report Date: 29-Sep-2004 08:40

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 29-SEP-2004 07:50
Lab File ID: UXJ24189.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m

COMPOUND	RRF	RF50	MIN	%D	MAX
41 Benzene	1.15116	1.16075 0.010	0.8	50.0	
42 Trichloroethene	0.26272	0.24593 0.010	-6.4	50.0	
43 1,2-Dichloropropane	0.28397	0.27945 0.010	-1.6	20.0	
44 1,4-Dioxane	0.00243	0.00283 0.010	16.3	50.0 <-	
45 Dibromomethane	0.15736	0.16442 0.010	4.5	50.0	
46 Bromodichloromethane	0.37355	0.36759 0.010	-1.6	50.0	
47 2-Chloroethyl vinyl ether	0.15967	0.15172 0.010	-5.0	50.0	
48 cis-1,3-Dichloropropene	0.45706	0.41930 0.010	-8.3	50.0	
49 4-Methyl-2-pentanone	0.24577	0.27896 0.010	13.5	50.0	
50 Toluene	1.49327	1.47633 0.010	-1.1	20.0	
51 trans-1,3-Dichloropropene	0.53539	0.46693 0.010	-12.8	50.0	
52 Ethyl Methacrylate	0.46066	0.46287 0.010	0.5	50.0	
53 1,1,2-Trichloroethane	0.30290	0.32224 0.010	6.4	50.0	
54 1,3-Dichloropropane	0.57071	0.56416 0.010	-1.1	50.0	
55 Tetrachloroethene	0.24193	0.24743 0.010	2.3	50.0	
56 2-Hexanone	0.24179	0.23807 0.010	-1.5	50.0	
57 Dibromochloromethane	0.32307	0.32805 0.010	1.5	50.0	
58 1,2-Dibromoethane	0.29844	0.29796 0.010	-0.2	50.0	
59 Chlorobenzene	0.96688	1.00040 0.300	3.5	50.0	
60 1,1,1,2-Tetrachloroethane	0.33590	0.33420 0.010	-0.5	50.0	
61 Ethylbenzene	0.48827	0.50140 0.010	2.7	20.0	
62 m + p-Xylene	0.62888	0.67860 0.010	7.9	50.0	
M 63 Xylenes (total)	0.62505	0.67651 0.010	8.2	50.0	
64 Xylene-o	0.61738	0.67233 0.010	8.9	50.0	
65 Styrene	1.10404	1.24068 0.010	12.4	50.0	
66 Bromoform	0.21960	0.22987 0.100	4.7	50.0	
67 Isopropylbenzene	1.37503	1.41026 0.010	2.6	50.0	
68 1,1,2,2-Tetrachloroethane	0.75371	0.79093 0.300	4.9	50.0	
69 1,4-Dichloro-2-butene	0.24799	0.23701 0.010	-4.4	50.0	
70 1,2,3-Trichloropropane	0.24733	0.25257 0.010	2.1	50.0	
71 Bromobenzene	0.74599	0.77267 0.010	3.6	50.0	
72 n-Propylbenzene	0.69551	0.74851 0.010	7.6	50.0	
73 2-Chlorotoluene	0.68879	0.70177 0.010	1.9	50.0	
74 1,3,5-Trimethylbenzene	2.31439	2.38191 0.010	2.9	50.0	
75 4-Chlorotoluene	0.73580	0.78061 0.010	6.1	50.0	
76 tert-Butylbenzene	1.87499	1.78120 0.010	-5.0	50.0	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24189.D
Report Date: 29-Sep-2004 08:40

STL North Canton

CONTINUING CALIBRATION COMPOUNDS

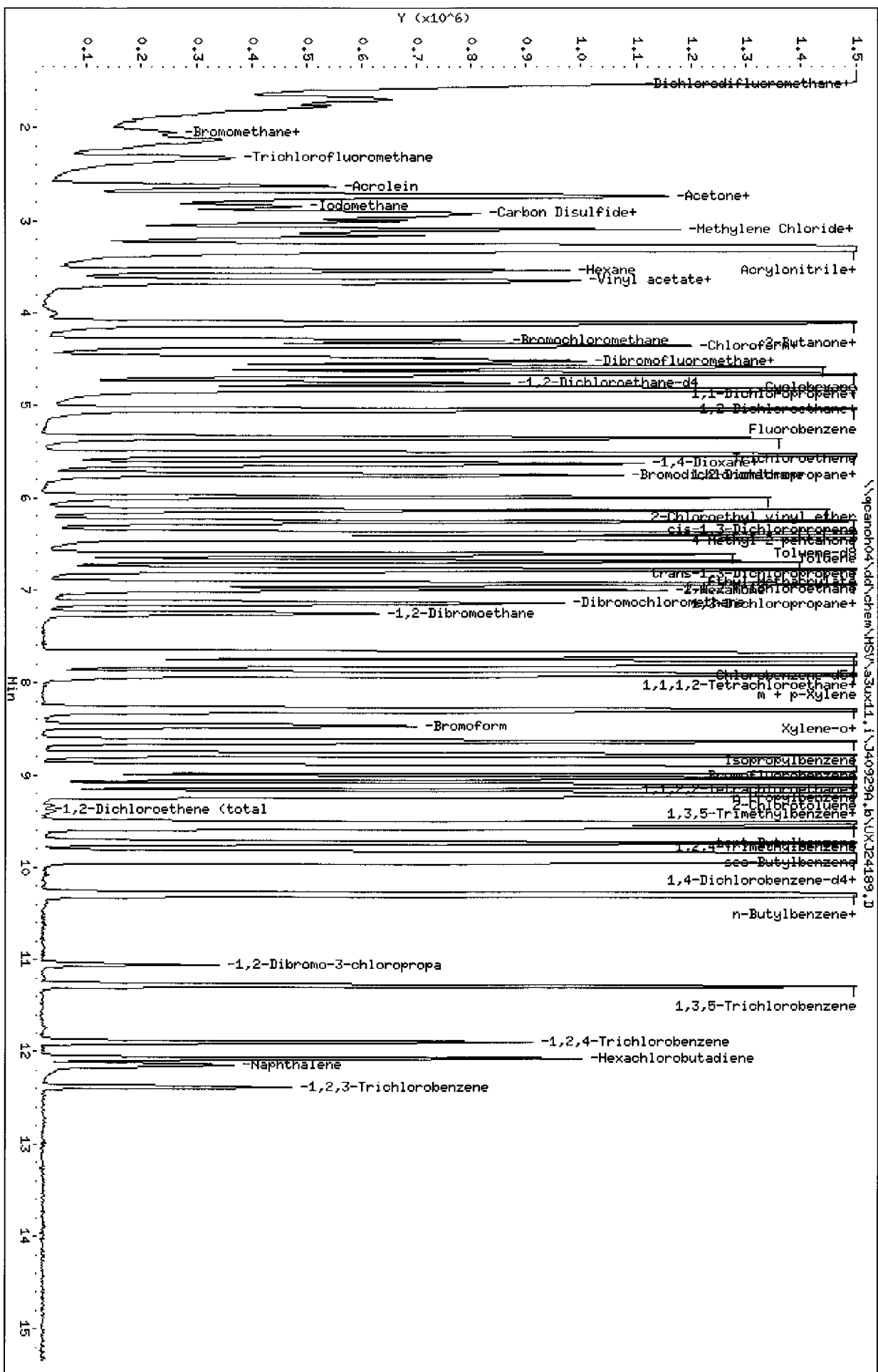
Instrument ID: a3ux11.i Injection Date: 29-SEP-2004 07:50
Lab File ID: UXJ24189.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m

COMPOUND	RRF	RF50	MIN	%D	MAX
77 1,2,4-Trimethylbenzene	2.47063	2.49994	0.010	1.2	50.0
78 sec-Butylbenzene	2.53792	2.45647	0.010	-3.2	50.0
79 4-Isopropyltoluene	2.11151	2.05723	0.010	-2.6	50.0
80 1,3-Dichlorobenzene	1.37723	1.36507	0.010	-0.9	50.0
81 1,4-Dichlorobenzene	1.44089	1.42117	0.010	-1.4	50.0
82 n-Butylbenzene	1.88626	1.71395	0.010	-9.1	50.0
83 1,2-Dichlorobenzene	1.35648	1.32540	0.010	-2.3	50.0
84 1,2-Dibromo-3-chloropropane	0.12763	0.09252	0.010	-27.5	50.0
85 1,2,4-Trichlorobenzene	0.57750	0.29334	0.010	-49.2	50.0
86 Hexachlorobutadiene	50.00000	41.15063	0.010	17.7	50.0
87 Naphthalene	1.36300	0.44117	0.010	-67.6	50.0 <-
88 1,2,3-Trichlorobenzene	0.41413	0.15677	0.010	-62.1	50.0 <-
98 Cyclohexane	0.36784	0.37366	0.010	1.6	50.0
143 Methyl Acetate	0.20568	0.19580	0.010	-4.8	50.0
144 Methylcyclohexane	0.30868	0.30279	0.010	-1.9	50.0
141 1,3,5-Trichlorobenzene	0.76231	0.60409	0.010	-20.8	50.0

Client ID:
Sample Info: 50NB-CC
Purge Volume: 5.0
Column phase: DB624

Instrument: ax3ux11.i

Operator: 43582
Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24189.D
Report Date: 30-Sep-2004 08:36

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24189.D
Lab Smp Id: 50NG-CC
Inj Date : 29-SEP-2004 07:50
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 50NG-CC
Misc Info : J40929A, 8260LLUX11, 2-8260.SUB, 43582, 2
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 1 Continuing Calibration Sample
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	ON-COL
*	1 Fluorobenzene	96	5.041	5.041 (1.000)	2187253	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1787660	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	1037377	50.0000		
\$	4 Dibromofluoromethane	113	4.473	4.473 (0.887)	534290	50.0000	52.466	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	742485	50.0000	53.022	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	2168826	50.0000	50.500	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	973862	50.0000	53.369	
8	Dichlorodifluoromethane	85	1.538	1.538 (0.305)	782348	50.0000	68.417	
9	Chloromethane	50	1.692	1.692 (0.336)	1147739	50.0000	56.216	
10	Vinyl Chloride	62	1.775	1.775 (0.352)	737730	50.0000	52.824	
11	Bromomethane	94	2.047	2.047 (0.406)	260299	50.0000	39.295	
12	Chloroethane	64	2.130	2.130 (0.423)	588879	50.0000	57.374	
13	Trichlorofluoromethane	101	2.319	2.319 (0.460)	620189	50.0000	42.368	
15	Acrolein	56	2.627	2.627 (0.521)	798155	500.000	584.18	
16	Acetone	43	2.734	2.734 (0.542)	418266	100.000	84.848	
17	1,1-Dichloroethene	96	2.722	2.722 (0.540)	493055	50.0000	50.407	
18	Freon-113	151	2.745	2.745 (0.545)	376435	50.0000	61.138	

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
19 Iodomethane	142	2.840	2.840 (0.563)		648021	50.0000	44.822
20 Carbon Disulfide	76	2.911	2.911 (0.578)		1930301	50.0000	52.830
21 Methylene Chloride	84	3.089	3.089 (0.613)		862596	50.0000	64.781
22 Acetonitrile	41	2.947	2.947 (0.585)		825584	500.000	635.98
23 Acrylonitrile	53	3.266	3.266 (0.648)		2188796	500.000	542.21
24 Methyl tert-butyl ether	73	3.313	3.313 (0.657)		1348723	50.0000	43.009
25 trans-1,2-Dichloroethene	96	3.313	3.313 (0.657)		578060	50.0000	49.817
26 Hexane	86	3.538	3.538 (0.702)		98487	50.0000	52.352
27 Vinyl acetate	43	3.668	3.668 (0.728)		849069	50.0000	45.728
28 1,1-Dichloroethane	63	3.645	3.645 (0.723)		1050947	50.0000	49.648
29 tert-Butyl Alcohol	59	3.160	3.160 (0.627)		750895	1000.00	882.63
30 2-Butanone	43	4.094	4.094 (0.812)		495988	100.000	82.791
M 31 1,2-Dichloroethene (total)	96				1150271	100.000	96.750
32 cis-1,2-dichloroethene	96	4.106	4.106 (0.815)		572211	50.0000	46.934
33 2,2-Dichloropropane	77	4.106	4.106 (0.815)		474981	50.0000	37.235
34 Bromochloromethane	128	4.296	4.296 (0.852)		292405	50.0000	51.794
35 Chloroform	83	4.355	4.355 (0.864)		1041680	50.0000	48.953
36 Tetrahydrofuran	42	4.343	4.343 (0.862)		150831	50.0000	45.603
37 1,1,1-Trichloroethane	97	4.520	4.520 (0.897)		705703	50.0000	43.912
38 1,1-Dichloropropene	75	4.651	4.651 (0.923)		747915	50.0000	49.934
39 Carbon Tetrachloride	117	4.662	4.662 (0.925)		619033	50.0000	49.058
40 1,2-Dichloroethane	62	4.816	4.816 (0.955)		862223	50.0000	49.592
41 Benzene	78	4.816	4.816 (0.955)		2538859	50.0000	50.417
42 Trichloroethene	130	5.349	5.349 (1.061)		537908	50.0000	46.804
43 1,2-Dichloropropane	63	5.526	5.526 (1.096)		611220	50.0000	49.204
44 1,4-Dioxane	88	5.633	5.633 (1.117)		309620	2500.00	2907.8 (A)
45 Dibromomethane	93	5.621	5.621 (1.115)		359621	50.0000	52.244
46 Bromodichloromethane	83	5.751	5.751 (1.141)		804008	50.0000	49.201
47 2-Chloroethyl vinyl ether	63	5.988	5.988 (1.188)		663705	100.000	95.022
48 cis-1,3-Dichloropropene	75	6.130	6.130 (1.216)		917124	50.0000	45.870
49 4-Methyl-2-pentanone	43	6.248	6.248 (1.239)		1220318	100.000	113.51
50 Toluene	91	6.437	6.437 (0.838)		2639185	50.0000	49.433
51 trans-1,3-Dichloropropene	75	6.603	6.603 (0.860)		834707	50.0000	43.606
52 Ethyl Methacrylate	69	6.686	6.686 (0.871)		827449	50.0000	50.240
53 1,1,2-Trichloroethane	97	6.769	6.769 (0.881)		576049	50.0000	53.193
54 1,3-Dichloropropane	76	6.922	6.922 (0.901)		1008519	50.0000	49.426
55 Tetrachloroethene	164	6.934	6.934 (0.903)		442315	50.0000	51.136
56 2-Hexanone	43	6.993	6.993 (0.911)		851161	100.000	98.458
57 Dibromochloromethane	129	7.135	7.135 (0.929)		586438	50.0000	50.770
58 1,2-Dibromoethane	107	7.254	7.254 (0.945)		532656	50.0000	49.920
59 Chlorobenzene	112	7.703	7.703 (1.003)		1788379	50.0000	51.734
60 1,1,1,2-Tetrachloroethane	131	7.774	7.774 (1.012)		597441	50.0000	49.747
61 Ethylbenzene	106	7.798	7.798 (1.015)		896339	50.0000	51.345
62 m + p-Xylene	106	7.905	7.905 (1.029)		2426225	100.000	107.91
M 63 Xylenes (total)	106				3628131	150.000	162.36
64 Xylene-o	106	8.283	8.283 (1.079)		1201906	50.0000	54.450
65 Styrene	104	8.295	8.295 (1.080)		2217918	50.0000	56.188

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
66 Bromoform	173	8.473	8.473 (1.103)	410935	50.0000	52.340	
67 Isopropylbenzene	105	8.626	8.626 (1.123)	2521070	50.0000	51.281	
68 1,1,2,2-Tetrachloroethane	83	8.899	8.899 (0.898)	820494	50.0000	52.469	
69 1,4-Dichloro-2-butene	53	8.958	8.958 (0.904)	245871	50.0000	47.787	
70 1,2,3-Trichloropropane	110	8.946	8.946 (0.903)	262012	50.0000	51.059	
71 Bromobenzene	156	8.934	8.934 (0.902)	801555	50.0000	51.788	
72 n-Propylbenzene	120	9.029	9.029 (0.912)	776488	50.0000	53.810	
73 2-Chlorotoluene	126	9.112	9.112 (0.920)	727999	50.0000	50.942	
74 1,3,5-Trimethylbenzene	105	9.194	9.194 (0.928)	2470939	50.0000	51.459	
75 4-Chlorotoluene	126	9.218	9.218 (0.931)	809789	50.0000	53.045	
76 tert-Butylbenzene	119	9.514	9.514 (0.961)	1847777	50.0000	47.499	
77 1,2,4-Trimethylbenzene	105	9.561	9.561 (0.965)	2593382	50.0000	50.593	
78 sec-Butylbenzene	105	9.727	9.727 (0.982)	2548287	50.0000	48.395	
79 4-Isopropyltoluene	119	9.869	9.869 (0.996)	2134120	50.0000	48.715	
80 1,3-Dichlorobenzene	146	9.845	9.845 (0.994)	1416093	50.0000	49.559	
81 1,4-Dichlorobenzene	146	9.928	9.928 (1.002)	1474292	50.0000	49.316	
82 n-Butylbenzene	91	10.271	10.271 (1.037)	1778009	50.0000	45.432	
83 1,2-Dichlorobenzene	146	10.295	10.295 (1.039)	1374938	50.0000	48.854	
84 1,2-Dibromo-3-chloropropane	157	11.052	11.052 (1.116)	95976	50.0000	36.244	
85 1,2,4-Trichlorobenzene	180	11.892	11.892 (1.201)	304304	50.0000	25.397	
86 Hexachlorobutadiene	225	12.070	12.070 (1.219)	219646	50.0000	41.151	
87 Naphthalene	128	12.141	12.141 (1.226)	457657	50.0000	16.184	
88 1,2,3-Trichlorobenzene	180	12.377	12.377 (1.250)	162634	50.0000	18.928	
98 Cyclohexane	56	4.580	4.580 (0.908)	817299	50.0000	50.792	
143 Methyl Acetate	43	3.006	3.006 (0.596)	856543	100.000	95.199	
144 Methylcyclohexane	83	5.514	5.514 (1.094)	662273	50.0000	49.046	
141 1,3,5-Trichlorobenzene	180	11.277	11.277 (1.139)	626668	50.0000	39.622	

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

Data File: \\QCANOH04\dd\chem\MSV\A3UX11.I\J40929A.D/UXJ24190.D
Report Date: 09/29/2004

CONTINUING CALIBRATION COMPOUNDS
PERCENT DRIFT REPORT

Instrument ID: a3ux11.i
Lab File ID: UXJ24190.D
Analysis Type: WATER

Injection Date: 29-SEP-2004 08:14
Lab Sample ID: 50NG-A9CC
Method File: \\QCANOH04\dd\chem\MSV\A3UX11.I\J40929A

COMPOUND	EXPECTED	MEASURED	%D	%D	MAX
	CONC.	CONC.			
39 Chlorobenzene-d5	50.0000	50.0000	0.0	50.0	
53 3-Chloropropene	50.0000	63.6847	27.4	50.0	
54 2-Chloro-1,3-butadiene	50.0000	53.0422	6.1	50.0	
55 Propionitrile	100.0000	100.2771	0.3	50.0	
56 Methacrylonitrile	50.0000	53.0943	6.2	50.0	
57 Isobutanol	1000.0000	1038.7096	3.9	50.0	
58 Methyl Methacrylate	50.0000	55.1079	10.2	50.0	
73 n-Butanol	1000.0000	986.9835	1.3	50.0	
74 Ethyl Acetate	100.0000	108.3418	8.3	50.0	
75 Cyclohexanone	500.0000	424.6793	15.1	50.0	
76 Ethyl Ether	50.0000	52.0369	4.1	50.0	
85 Dichlorofluoromethane	50.0000	64.2558	28.5	50.0	
86 2-Nitropropane	100.0000	103.2894	3.3	50.0	
126 Isopropyl Ether	250.0000	259.5683	3.8	50.0	
130 Fluorobenzene	50.0000	50.0000	0.0	50.0	
132 1,4-Dichlorobenzene-d4	50.0000	50.0000	0.0	50.0	

Data File: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24190.D
Report Date: 29-Sep-2004 08:29

STL - North Canton

CONTINUING CALIBRATION COMPOUNDS

Instrument ID: a3ux11.i Injection Date: 29-SEP-2004 08:14
Lab File ID: UXJ24190.D Init. Cal. Date(s): 16-AUG-2004 14-SEP-2004
Analysis Type: WATER Init. Cal. Times: 16:18 15:41
Lab Sample ID: 50NG-A9CC Quant Type: ISTD
Method: \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m

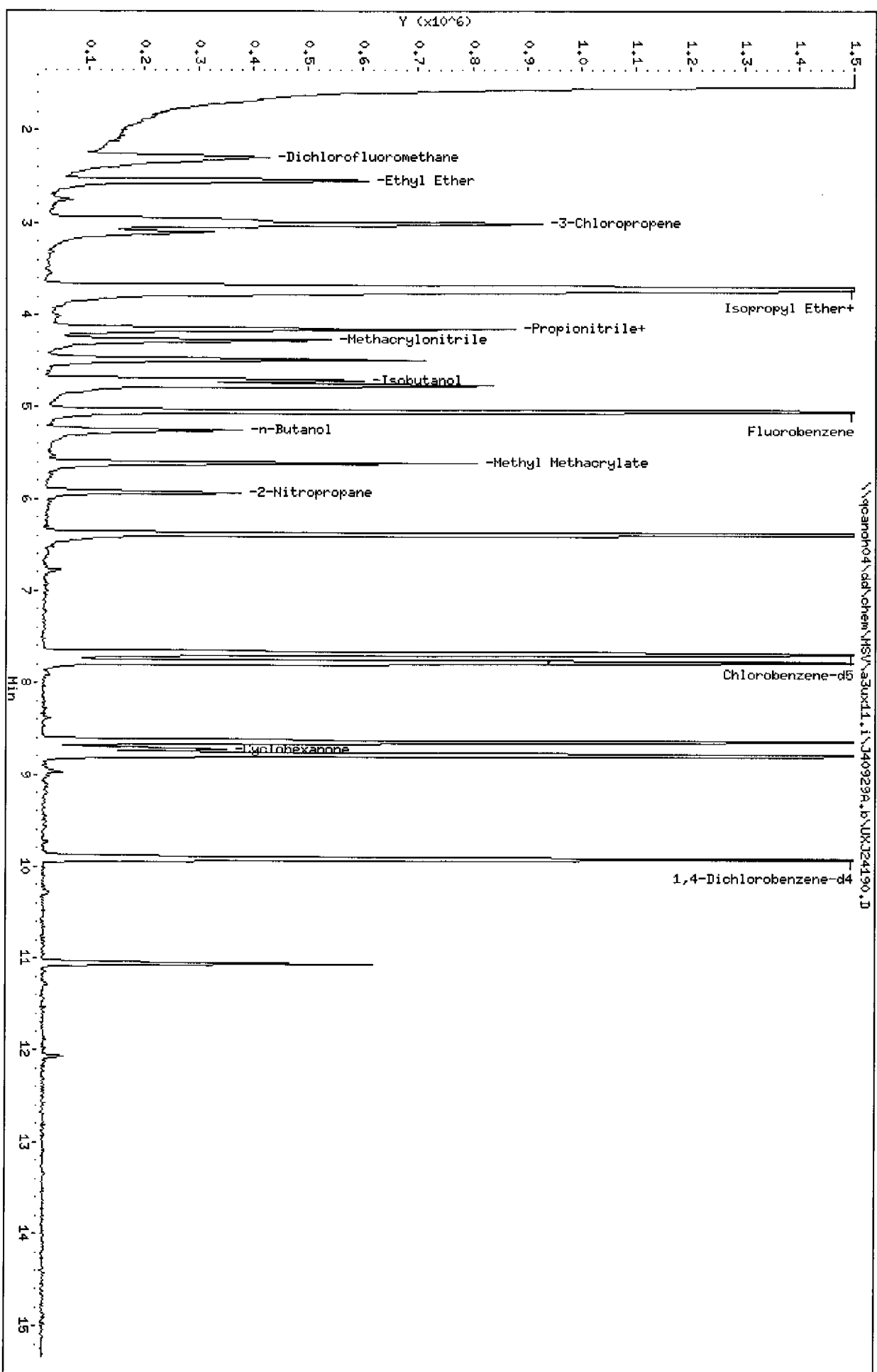
COMPOUND	RRF	RF50	MIN	%D	MAX
14 Dichlorofluoromethane	0.48495	0.62322	0.010	28.5	50.0
89 Ethyl Ether	0.24654	0.25659	0.010	4.1	50.0
91 3-Chloropropene	0.10305	0.13125	0.010	27.4	50.0
92 Isopropyl Ether	0.22353	0.23209	0.010	3.8	50.0
93 2-Chloro-1,3-butadiene	0.37276	0.39544	0.010	6.1	50.0
94 Propionitrile	0.04231	0.04243	0.010	0.3	50.0
95 Ethyl Acetate	0.24508	0.26553	0.010	8.3	50.0
96 Methacrylonitrile	0.15890	0.16873	0.010	6.2	50.0
97 Isobutanol	0.01142	0.01186	0.010	3.9	50.0
99 n-Butanol	0.00822	0.00811	0.010	-1.3	50.0
100 Methyl Methacrylate	0.19531	0.21526	0.010	10.2	50.0
101 2-Nitropropane	0.06079	0.06279	0.010	3.3	50.0
103 Cyclohexanone	0.02717	0.02308	0.010	-15.1	50.0

Data File: \\qcarnoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UKJ24190.D
Date : 29-SEP-2004 08:14
Client ID:
Sample Info: E01C-A9CC

Purge Volume: 5.0
Column Phase: DB624

Instrument: a3ux11.i
Operator: 43582
Column diameter: 0.18

\\qcarnoh04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UKJ24190.D



Data File: \\qcanoh04\dd\chem\MSV\A3UX11.i\J40929A.b\UXJ24190.D
Report Date: 29-Sep-2004 08:39

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\A3UX11.i\J40929A.b\UXJ24190.D
Lab Smp Id: 50NG-A9CC
Inj Date : 29-SEP-2004 08:14
Operator : 43582 Inst ID: a3ux11.i
Smp Info : 50NG-A9CC
Misc Info : J40929A,8260LLUX11,3-IX.SUB,43582,2
Comment :
Method : \\QCANOH04\dd\chem\MSV\A3UX11.i\J40929A.b\8260LLUX11.m
Meth Date : 29-Sep-2004 08:39 evans1 Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 2 Continuing Calibration Sample
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 3-IX.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	AMOUNTS					
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ng)
* 1 Fluorobenzene	96	5.041	5.041 (1.000)	2085210	50.0000		
* 2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1694750	50.0000		
* 3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	774505	50.0000		
14 Dichlorofluoromethane	67	2.284	2.284 (0.453)	1299534	50.0000	64.256	
89 Ethyl Ether	59	2.544	2.544 (0.505)	535037	50.0000	52.037	
91 3-Chloropropene	76	3.018	3.018 (0.599)	273692	50.0000	63.684	
92 Isopropyl Ether	87	3.704	3.704 (0.735)	2419748	250.000	259.57 (A)	
93 2-Chloro-1,3-butadiene	53	3.728	3.728 (0.739)	824568	50.0000	53.042	
94 Propionitrile	54	4.142	4.142 (0.822)	176932	100.000	100.28	
95 Ethyl Acetate	43	4.154	4.154 (0.824)	1107359	100.000	108.34	
96 Methacrylonitrile	41	4.272	4.272 (0.847)	351838	50.0000	53.094	
97 Isobutanol	41	4.710	4.710 (0.613)	402003	1000.00	1038.7 (A)	
99 n-Butanol	56	5.254	5.254 (0.684)	274928	1000.00	986.98 (A)	
100 Methyl Methacrylate	41	5.609	5.609 (1.113)	448862	50.0000	55.108	
101 2-Nitropropane	41	5.928	5.928 (1.176)	261879	100.000	103.29	
103 Cyclohexanone	55	8.721	8.721 (0.881)	178750	500.000	424.68 (A)	

Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24190.D
Report Date: 29-Sep-2004 08:39

QC Flag Legend

A - Target compound detected but, quantitated amount exceeded maximum amount.

RAW QC DATA

Date : 16-AUG-2004 13:09

Client ID: 50NG BFB

Instrument: z3ux11.i

Sample Info:

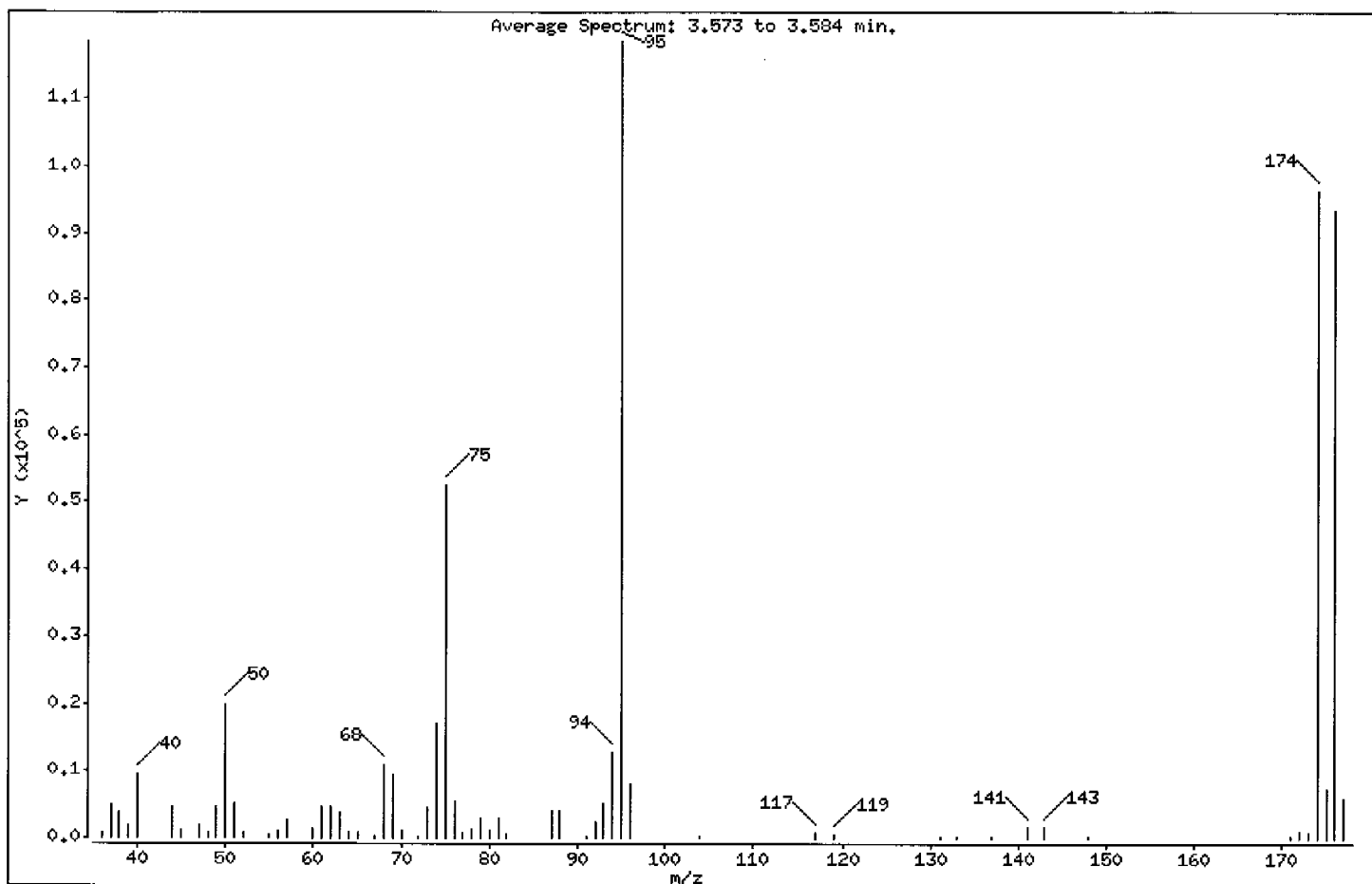
Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

1 kfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE	
95	Base Peak, 100% relative abundance	100.00	
50	15.00 - 40.00% of mass 95	16.76	
75	30.00 - 60.00% of mass 95	44.47	
96	5.00 - 9.00% of mass 95	6.93	
173	Less than 2.00% of mass 174	0.68 (< 0.84)	
174	50.00 - 100.00% of mass 95	81.27	
176	5.00 - 9.00% of mass 174	6.13 (< 7.55)	
176	95.00 - 101.00% of mass 174	78.91 (< 97.09)	
177	5.00 - 9.00% of mass 176	5.04 (< 6.39)	

Date : 16-AUG-2004 13:09

Client ID: 50NG BFB

Instrument: z3ux11.i

Sample Info:

Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

Data File: BFB207.D

Spectrum: Average Spectrum; 3.573 to 3.584 min.

Location of Maximum: 95.00

Number of points: 61

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	749	60.00	1376	78.00	1307	131.00	285
37.00	4867	61.00	4538	79.00	3095	133.00	271
38.00	3795	62.00	4718	80.00	1085	137.00	323
39.00	1854	63.00	3709	81.00	3055	141.00	1521
40.00	9546	64.00	700	82.00	515	143.00	1655
44.00	4571	65.00	821	87.00	4063	148.00	297
45.00	1190	67.00	254	88.00	4069	171.00	316
47.00	2040	68.00	10867	91.00	257	172.00	1062
48.00	737	69.00	9493	92.00	2338	173.00	806
49.00	4592	70.00	1139	93.00	5215	174.00	96288
50.00	19856	72.00	389	94.00	12726	175.00	7267
51.00	5302	73.00	4639	95.00	118480	176.00	93488
52.00	840	74.00	17288	96.00	8207	177.00	5974
55.00	613	75.00	52680	104.00	355		
56.00	1116	76.00	5372	117.00	715		
57.00	2804	77.00	828	119.00	414		

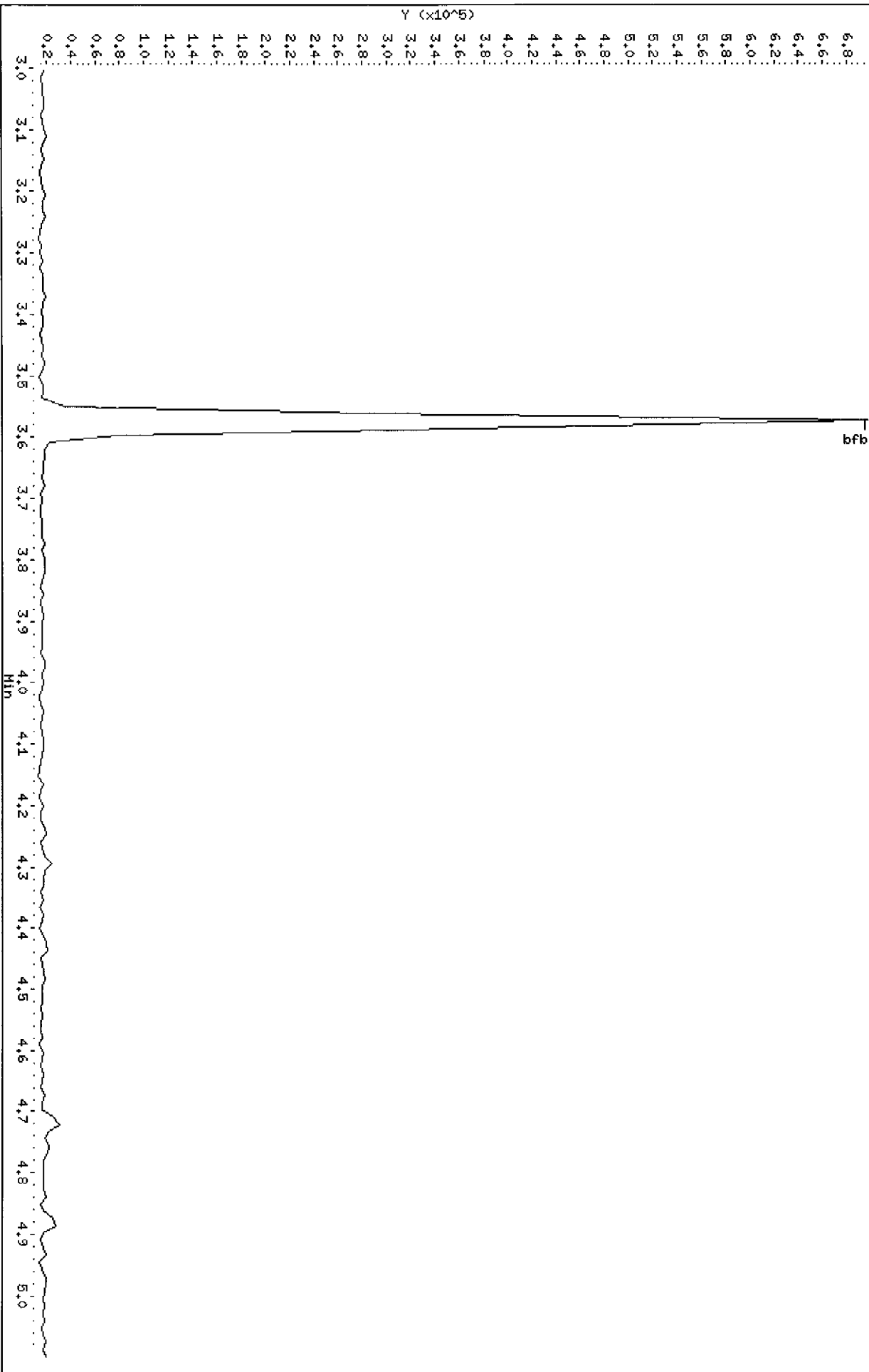
Client ID: 5ONG BEB

Sample Info:

Volume Injected (uL): 1.0
Column phase: DB624 20M

Instrument: a3ux11.i
Operator: 43582
Column diameter: 0.18

\\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40816A-IC.b\\BFB207.D



Data File: \\qcanch04\dd\chem\MSV\z3ux11.i\J40914B-1C.b\BFB232.D

Date : 14-SEP-2004 13:21

Client ID: 50NG BFB

Instrument: z3ux11.i

Sample Info: BFB232

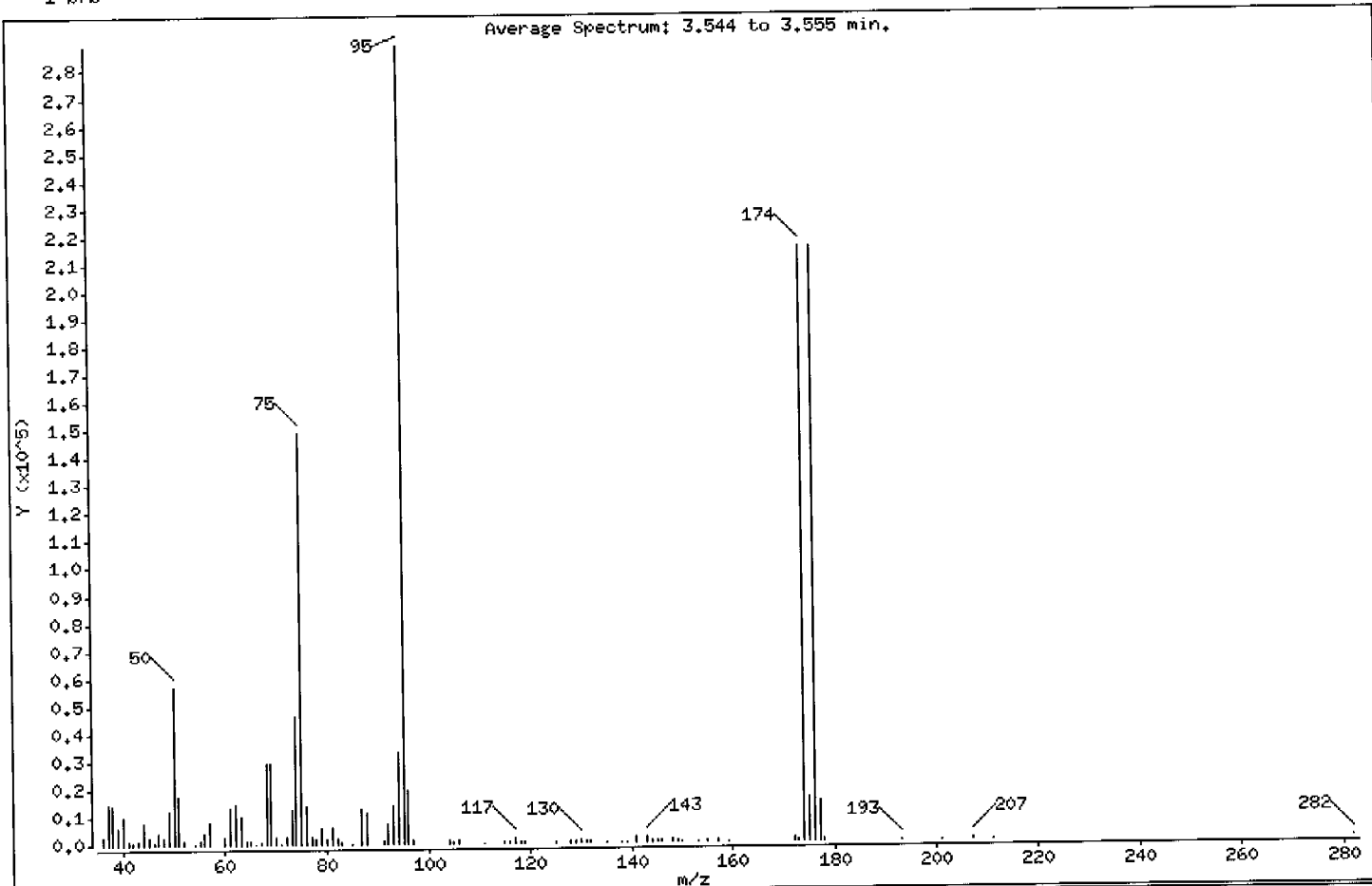
Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

1 kfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE	
95	Base Peak, 100% relative abundance	100.00	
50	15.00 - 40.00% of mass 95	19.83	
75	30.00 - 60.00% of mass 95	51.55	
96	5.00 - 9.00% of mass 95	6.64	
173	Less than 2.00% of mass 174	0.21 (< 0.28)	
174	50.00 - 100.00% of mass 95	74.58	
175	5.00 - 9.00% of mass 174	5.61 (< 7.52)	
176	95.00 - 101.00% of mass 174	74.49 (< 99.87)	
177	5.00 - 9.00% of mass 176	5.17 (< 6.94)	

Date : 14-SEP-2004 13:21

Client ID: 5ONG BFB

Instrument: z3ux11.i

Sample Info: BFB232

Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

Data File: BFB232.D

Spectrum: Average Spectrum: 3.544 to 3.555 min.

Location of Maximum: 35.00

Number of points: 97

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	2406	64.00	1324	93.00	13655	144.00	401
37.00	14778	65.00	1522	94.00	33528	145.00	409
38.00	13742	66.00	263	95.00	288960	146.00	877
39.00	5795	67.00	910	96.00	19192	148.00	1075
40.00	10147	68.00	29456	97.00	1451	149.00	479
41.00	1576	69.00	29344	104.00	1533	150.00	313
42.00	737	70.00	2634	105.00	833	153.00	287
43.00	1309	71.00	324	106.00	1128	155.00	668
44.00	8117	72.00	2356	111.00	260	157.00	711
45.00	2441	73.00	12818	115.00	375	159.00	271
46.00	749	74.00	46784	116.00	766	172.00	1628
47.00	3975	75.00	148928	117.00	1678	173.00	603
48.00	2388	76.00	13979	118.00	426	174.00	215488
49.00	12238	77.00	2377	119.00	915	175.00	16212
50.00	57288	78.00	2058	125.00	649	176.00	215232
51.00	17136	79.00	5934	128.00	783	177.00	14940
52.00	1094	80.00	2304	129.00	689	178.00	355
54.00	314	81.00	6191	130.00	1263	193.00	286
55.00	1279	82.00	1726	131.00	957	201.00	259
56.00	4132	83.00	677	132.00	360	207.00	652
57.00	7889	85.00	327	135.00	330	211.00	272
60.00	2651	87.00	12709	138.00	255	282.00	251
61.00	13040	88.00	11543	139.00	322		
62.00	14627	91.00	1380	141.00	2064		
63.00	10230	92.00	7594	143.00	2214		

Data File: \\pcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\BF232.D

Date : 14-SEP-2004 13:21

Client ID: SONG-BFB

Sample Info: BF232

Volume Injected (uL) : 1.0

Column phase: DB624 20M

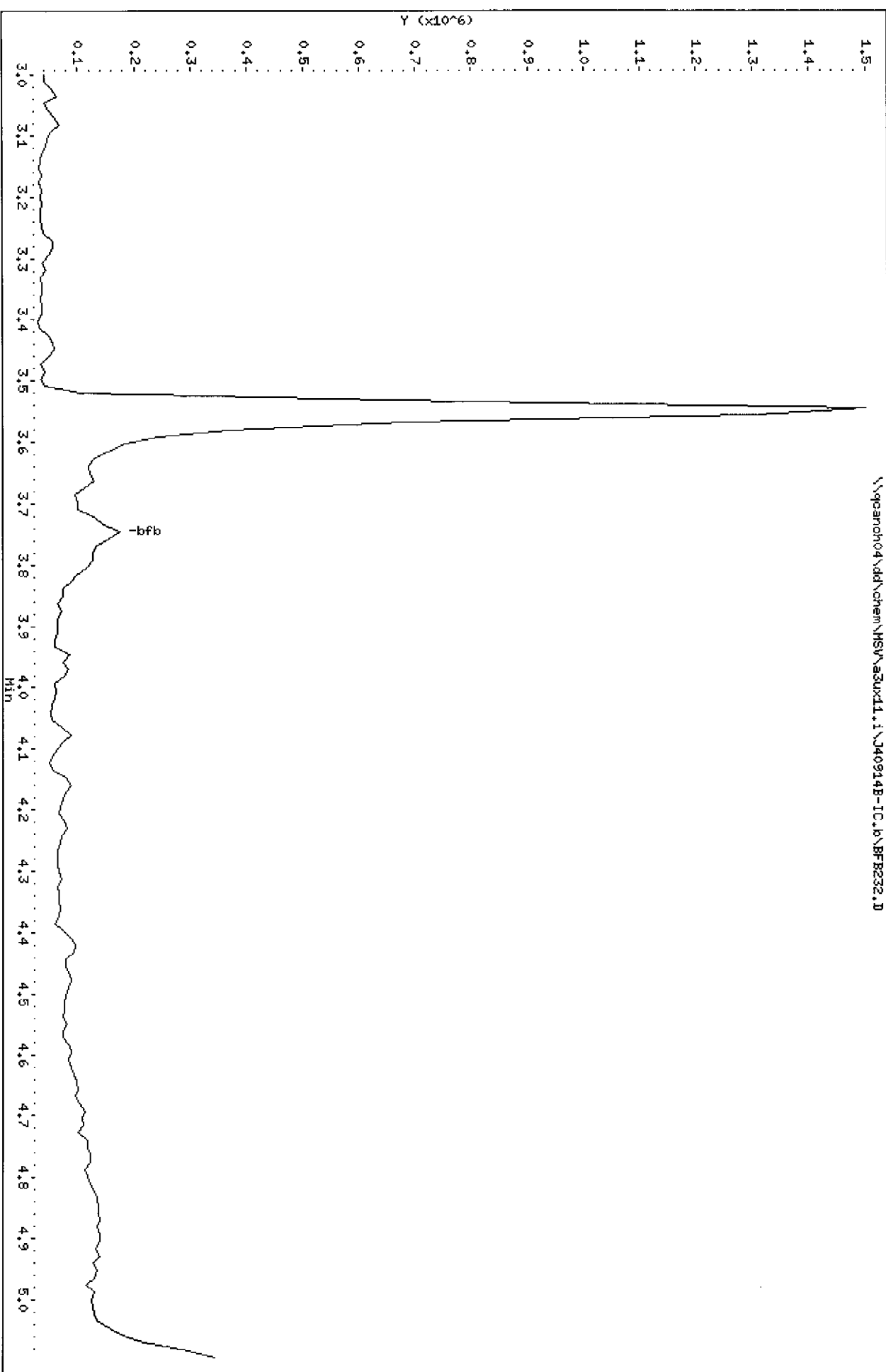
Page 2

Instrument: a3ux11.i

Operator: 43582

Column diameter: 0.18

\\pcanoh04\\dd\\chem\\MSV\\a3ux11.i\\J40914B-IC.b\\BF232.D



Date : 29-SEP-2004 07:25

Client ID: SONG BFB

Instrument: z3ux11.i

Sample Info:

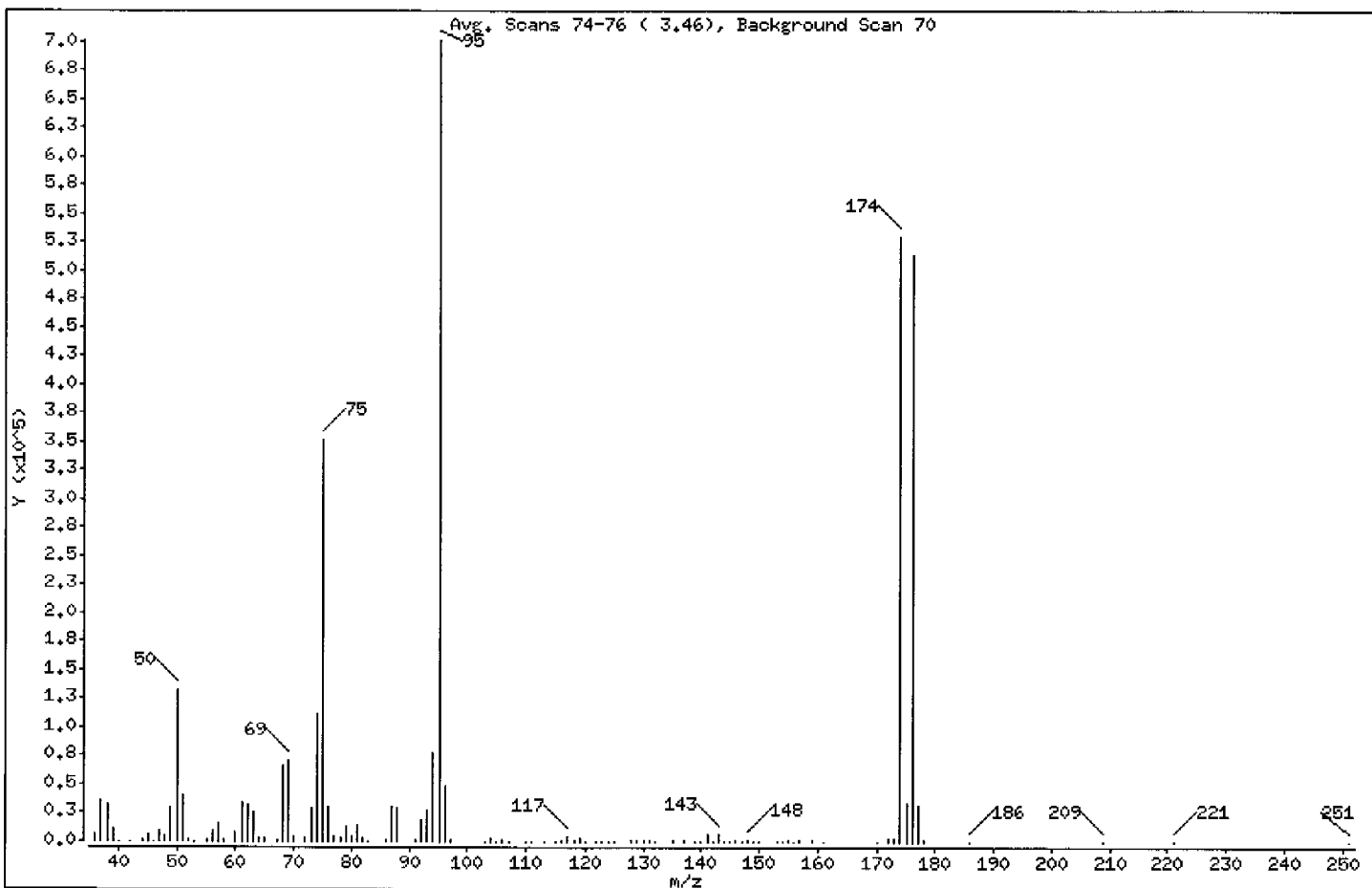
Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95 Base Peak, 100% relative abundance	100,00	
50 15.00 - 40.00% of mass 95	18,79	
75 30.00 - 60.00% of mass 95	50,16	
96 5.00 - 9.00% of mass 95	6,82	
173 Less than 2.00% of mass 174	0,41 (< 0,54)	
174 50,00 - 100,00% of mass 95	75,53	
175 5.00 - 9.00% of mass 174	4,87 (< 6,45)	
176 95,00 - 101,00% of mass 174	73,31 (< 97,06)	
177 5.00 - 9.00% of mass 176	4,63 (< 6,32)	

Date : 29-SEP-2004 07:25

Client ID: SONG BFB

Instrument: c3ux11.i

Sample Info:

Volume Injected (uL): 1.0

Operator: 43582

Column phase: DB624 20M

Column diameter: 0.18

Data File: BFB243.D

Spectrum: Avg. Scans 74-76 (3,46), Background Scan 70

Location of Maximum: 95.00

Number of points: 108

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	5960	69.00	70248	107.00	770	146.00	1100
37.00	35008	70.00	4668	110.00	183	147.00	384
38.00	32560	72.00	3224	111.00	467	148.00	1809
39.00	11582	73.00	29128	113.00	202	149.00	448
40.00	543	74.00	111896	115.00	751	150.00	766
42.00	400	75.00	351552	116.00	2358	153.00	297
44.00	1687	76.00	30440	117.00	4467	154.00	261
45.00	5920	77.00	4123	118.00	2017	155.00	1349
46.00	384	78.00	2841	119.00	2626	156.00	251
47.00	10446	79.00	13102	120.00	171	157.00	1102
48.00	4227	80.00	4752	122.00	204	159.00	917
49.00	28776	81.00	15207	123.00	253	161.00	524
50.00	131648	82.00	3025	124.00	499	170.00	176
51.00	40632	83.00	437	125.00	189	172.00	3122
52.00	1144	86.00	907	128.00	2323	173.00	2884
53.00	287	87.00	30824	129.00	1074	174.00	529344
55.00	1721	88.00	28960	130.00	1788	175.00	34144
56.00	8900	91.00	1910	131.00	1154	176.00	513792
57.00	16624	92.00	19328	132.00	384	177.00	32448
58.00	1009	93.00	27192	135.00	1133	178.00	1312
60.00	7276	94.00	76880	137.00	1455	186.00	168
61.00	33232	95.00	700864	139.00	177	209.00	176
62.00	32496	96.00	47792	140.00	456	221.00	174
63.00	25144	97.00	1224	141.00	6397	251.00	470
64.00	3398	103.00	167	142.00	739		
65.00	3847	104.00	2568	143.00	6467		
67.00	2045	105.00	711	144.00	614		
68.00	66864	106.00	2074	145.00	747		

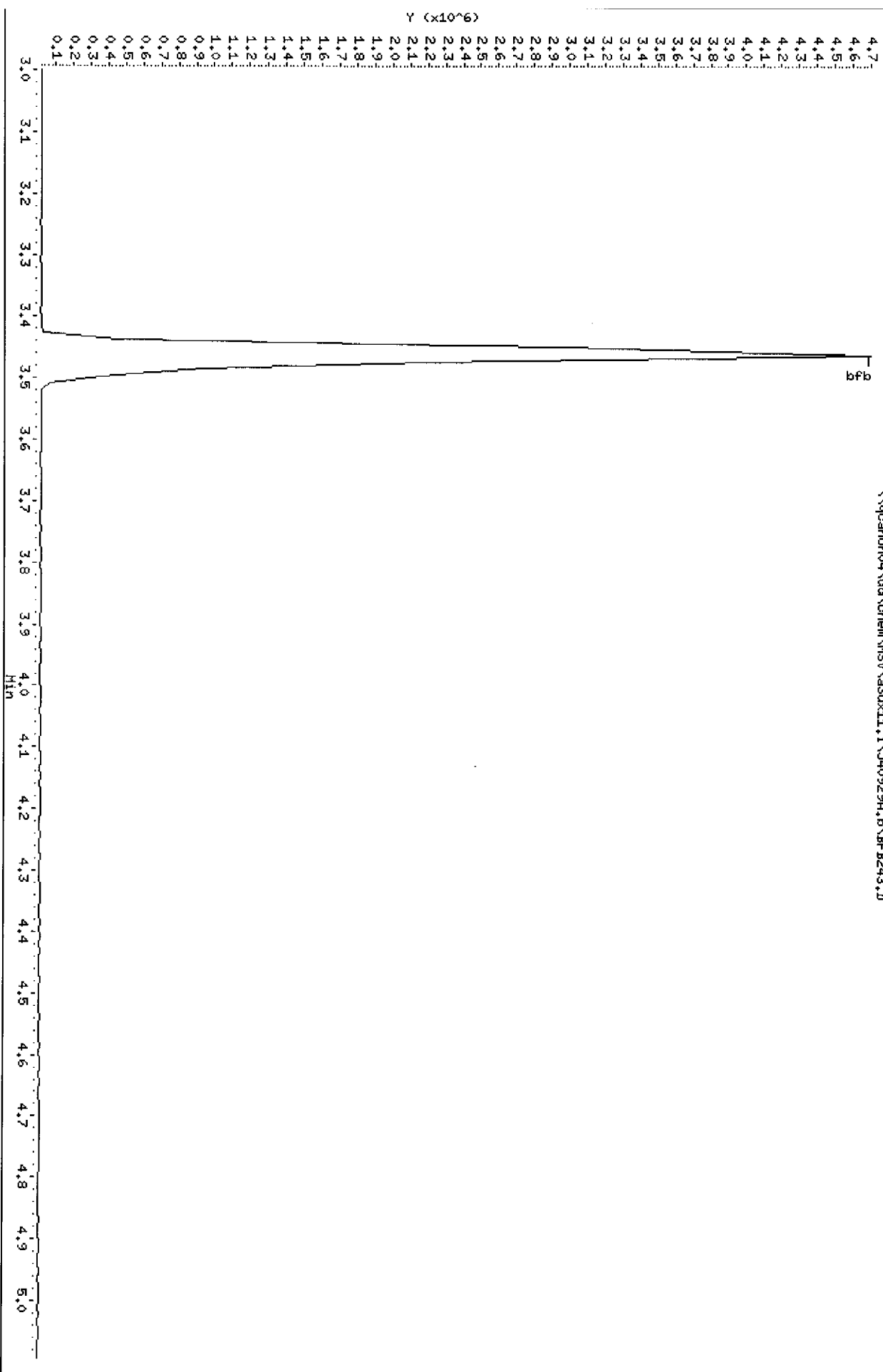
Client ID: 5ONG.BFB

Sample Info:

Volume Injected (uL): 1.0
Column Phase: DB624 20M

Instrument: a3ux11.i
Operator: 43582
Column diameter: 0.18

\\qcando04\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\BF243.D



LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

PARAMETER	PERCENT RECOVERY	RECOVERY LIMITS		RPD LIMITS	METHOD
		RPD	LIMITS		
Acetone	89	(22 - 200)			SW846 8260B
	91	(22 - 200)	2.1	(0-95)	SW846 8260B
Benzene	99	(80 - 116)			SW846 8260B
	97	(80 - 116)	1.9	(0-20)	SW846 8260B
Bromodichloromethane	101	(87 - 130)			SW846 8260B
	99	(87 - 130)	1.4	(0-30)	SW846 8260B
Bromoform	106	(76 - 150)			SW846 8260B
	110	(76 - 150)	3.5	(0-30)	SW846 8260B
Bromomethane	73	(64 - 129)			SW846 8260B
	78	(64 - 129)	6.6	(0-30)	SW846 8260B
2-Butanone	76	(28 - 237)			SW846 8260B
	87	(28 - 237)	12	(0-65)	SW846 8260B
Carbon disulfide	98	(73 - 139)			SW846 8260B
	104	(73 - 139)	5.5	(0-30)	SW846 8260B
Carbon tetrachloride	94	(75 - 149)			SW846 8260B
	96	(75 - 149)	1.4	(0-30)	SW846 8260B
Chlorobenzene	102	(76 - 117)			SW846 8260B
	100	(76 - 117)	2.6	(0-20)	SW846 8260B
Dichlorodifluoromethane	64 a	(70 - 130)			SW846 8260B
	69 a	(70 - 130)	7.1	(0-30)	SW846 8260B
1,1,2-Trichloro- 1,2,2-trifluoroethane	124	(70 - 130)			SW846 8260B
	126	(70 - 130)	1.7	(0-30)	SW846 8260B
Methyl acetate	95	(70 - 130)			SW846 8260B
	100	(70 - 130)	5.0	(0-30)	SW846 8260B
Methyl tert-butyl ether (MTBE)	86	(70 - 130)			SW846 8260B
	83	(70 - 130)	3.5	(0-30)	SW846 8260B
Cyclohexane	88	(70 - 130)			SW846 8260B
	92	(70 - 130)	4.0	(0-30)	SW846 8260B
Methylcyclohexane	87	(70 - 130)			SW846 8260B
	90	(70 - 130)	3.1	(0-30)	SW846 8260B
Dibromochloromethane	101	(81 - 138)			SW846 8260B
	99	(81 - 138)	1.3	(0-30)	SW846 8260B
Isopropylbenzene	110	(70 - 130)			SW846 8260B
	111	(70 - 130)	1.2	(0-30)	SW846 8260B

(Continued on next page)

LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

Client Lot #....: A4I270179 Work Order #....: GRDLR1AC-LCS Matrix.....: WATER
LCS Lot-Sample#: A4I290000-268 GRDLR1AD-LCSD

PARAMETER	PERCENT RECOVERY	RECOVERY LIMITS	RPD		METHOD
			LIMITS	RPD	
1,3-Dichlorobenzene	98	(70 - 130)			SW846 8260B
	94	(70 - 130)	4.5	(0-30)	SW846 8260B
Chloroethane	92	(66 - 126)			SW846 8260B
	96	(66 - 126)	4.6	(0-30)	SW846 8260B
1,4-Dichlorobenzene	103	(70 - 130)			SW846 8260B
	100	(70 - 130)	2.8	(0-30)	SW846 8260B
1,2-Dichlorobenzene	98	(70 - 130)			SW846 8260B
	94	(70 - 130)	3.6	(0-30)	SW846 8260B
1,2,4-Trichloro- benzene	36 a	(70 - 130)			SW846 8260B
	40 a	(70 - 130)	11	(0-30)	SW846 8260B
Chloroform	99	(84 - 128)			SW846 8260B
	97	(84 - 128)	2.4	(0-30)	SW846 8260B
Chloromethane	78	(48 - 123)			SW846 8260B
	82	(48 - 123)	4.1	(0-30)	SW846 8260B
1,2-Dibromo-3-chloro- propane	76	(70 - 130)			SW846 8260B
	78	(70 - 130)	2.2	(0-30)	SW846 8260B
1,2-Dibromoethane	98	(70 - 130)			SW846 8260B
	100	(70 - 130)	1.9	(0-30)	SW846 8260B
1,1-Dichloroethane	98	(86 - 123)			SW846 8260B
	100	(86 - 123)	1.8	(0-30)	SW846 8260B
1,2-Dichloroethane	101	(79 - 136)			SW846 8260B
	101	(79 - 136)	0.18	(0-30)	SW846 8260B
cis-1,2-Dichloroethene	94	(85 - 113)			SW846 8260B
	95	(85 - 113)	1.2	(0-30)	SW846 8260B
trans-1,2-Dichloroethene	97	(79 - 120)			SW846 8260B
	102	(79 - 120)	5.3	(0-30)	SW846 8260B
1,1-Dichloroethene	101	(63 - 130)			SW846 8260B
	103	(63 - 130)	1.8	(0-20)	SW846 8260B
1,2-Dichloroethene (total)	96	(82 - 116)			SW846 8260B
	99	(82 - 116)	3.3	(0-30)	SW846 8260B
1,2-Dichloropropane	100	(82 - 115)			SW846 8260B
	96	(82 - 115)	3.4	(0-30)	SW846 8260B
cis-1,3-Dichloropropene	92	(84 - 130)			SW846 8260B
	92	(84 - 130)	0.13	(0-30)	SW846 8260B
trans-1,3-Dichloropropene	85	(84 - 130)			SW846 8260B
	84	(84 - 130)	1.0	(0-30)	SW846 8260B

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LABORATORY CONTROL SAMPLE EVALUATION REPORT

GC/MS Volatiles

PARAMETER	PERCENT	RECOVERY	RPD	LIMITS	METHOD
	RECOVERY	LIMITS			
Ethylbenzene	100	(86 - 116)			SW846 8260B
	100	(86 - 116)	0.71	(0-30)	SW846 8260B
2-Hexanone	81	(35 - 200)			SW846 8260B
	86	(35 - 200)	6.0	(0-52)	SW846 8260B
Methylene chloride	133 a	(78 - 118)			SW846 8260B
	138 a	(78 - 118)	3.8	(0-30)	SW846 8260B
4-Methyl-2-pentanone	107	(78 - 141)			SW846 8260B
	117	(78 - 141)	8.9	(0-32)	SW846 8260B
Styrene	110	(85 - 117)			SW846 8260B
	108	(85 - 117)	1.8	(0-30)	SW846 8260B
1,1,2,2-Tetrachloroethane	113	(85 - 118)			SW846 8260B
	116	(85 - 118)	2.3	(0-30)	SW846 8260B
Tetrachloroethene	96	(88 - 113)			SW846 8260B
	98	(88 - 113)	1.6	(0-30)	SW846 8260B
Toluene	95	(74 - 119)			SW846 8260B
	94	(74 - 119)	1.2	(0-20)	SW846 8260B
1,1,1-Trichloroethane	88	(78 - 140)			SW846 8260B
	87	(78 - 140)	1.4	(0-30)	SW846 8260B
1,1,2-Trichloroethane	98	(83 - 122)			SW846 8260B
	99	(83 - 122)	1.3	(0-30)	SW846 8260B
Trichloroethene	96	(75 - 122)			SW846 8260B
	96	(75 - 122)	0.010	(0-20)	SW846 8260B
Trichlorofluoromethane	80	(70 - 130)			SW846 8260B
	82	(70 - 130)	1.8	(0-30)	SW846 8260B
Vinyl chloride	80	(61 - 120)			SW846 8260B
	82	(61 - 120)	2.5	(0-30)	SW846 8260B
Xylenes (total)	106	(87 - 116)			SW846 8260B
	105	(87 - 116)	0.96	(0-30)	SW846 8260B

<u>SURROGATE</u>	<u>PERCENT RECOVERY</u>	<u>RECOVERY LIMITS</u>
Dibromofluoromethane	104	(73 - 122)
1,2-Dichloroethane-d4	104	(73 - 122)
Toluene-d8	104	(61 - 128)
4-Bromofluorobenzene	98	(76 - 110)
	97	(76 - 110)
	108	(74 - 116)
	109	(74 - 116)

NOTE (S) :

Calculations are performed before rounding to avoid round-off errors in calculated results.

Bold print denotes control parameters

a Spiked analyte recovery is outside stated control limits

LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

PARAMETER	SPIKE	MEASURED		PERCENT		METHOD
	AMOUNT	AMOUNT	UNITS	RECOVERY	RPD	
Acetone	10	8.9	ug/L	89		SW846 8260B
	10	9.1	ug/L	91	2.1	SW846 8260B
Benzene	10	9.9	ug/L	99		SW846 8260B
	10	9.7	ug/L	97	1.9	SW846 8260B
Bromodichloromethane	10	10	ug/L	101		SW846 8260B
	10	9.9	ug/L	99	1.4	SW846 8260B
Bromoform	10	11	ug/L	106		SW846 8260B
	10	11	ug/L	110	3.5	SW846 8260B
Bromomethane	10	7.3	ug/L	73		SW846 8260B
	10	7.8	ug/L	78	6.6	SW846 8260B
2-Butanone	10	7.6	ug/L	76		SW846 8260B
	10	8.7	ug/L	87	12	SW846 8260B
Carbon disulfide	10	9.8	ug/L	98		SW846 8260B
	10	10	ug/L	104	5.5	SW846 8260B
Carbon tetrachloride	10	9.4	ug/L	94		SW846 8260B
	10	9.6	ug/L	96	1.4	SW846 8260B
Chlorobenzene	10	10	ug/L	102		SW846 8260B
	10	10	ug/L	100	2.6	SW846 8260B
Dichlorodifluoromethane	10	6.4 a	ug/L	64		SW846 8260B
	10	6.9 a	ug/L	69	7.1	SW846 8260B
1,1,2-Trichloro- 1,2,2-trifluoroethane	10	12	ug/L	124		SW846 8260B
	10	13	ug/L	126	1.7	SW846 8260B
Methyl acetate	10	9.5	ug/L	95		SW846 8260B
	10	10	ug/L	100	5.0	SW846 8260B
Methyl tert-butyl ether (MTBE)	10	8.6	ug/L	86		SW846 8260B
	10	8.3	ug/L	83	3.5	SW846 8260B
Cyclohexane	10	8.8	ug/L	88		SW846 8260B
	10	9.2	ug/L	92	4.0	SW846 8260B
Methylcyclohexane	10	8.7	ug/L	87		SW846 8260B
	10	9.0	ug/L	90	3.1	SW846 8260B
Dibromochloromethane	10	10	ug/L	101		SW846 8260B
	10	9.9	ug/L	99	1.3	SW846 8260B
Isopropylbenzene	10	11	ug/L	110		SW846 8260B
	10	11	ug/L	111	1.2	SW846 8260B

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LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

PARAMETER	SPIKE	MEASURED		PERCENT		METHOD
	AMOUNT	AMOUNT	UNITS	RECOVERY	RPD	
1,3-Dichlorobenzene	10	9.8	ug/L	98		SW846 8260B
	10	9.4	ug/L	94	4.5	SW846 8260B
Chloroethane	10	9.2	ug/L	92		SW846 8260B
	10	9.6	ug/L	96	4.6	SW846 8260B
1,4-Dichlorobenzene	10	10	ug/L	103		SW846 8260B
	10	10	ug/L	100	2.8	SW846 8260B
1,2-Dichlorobenzene	10	9.8	ug/L	98		SW846 8260B
	10	9.4	ug/L	94	3.6	SW846 8260B
1,2,4-Trichloro-benzene	10	3.6 a	ug/L	36		SW846 8260B
	10	4.0 a	ug/L	40	11	SW846 8260B
Chloroform	10	9.9	ug/L	99		SW846 8260B
	10	9.7	ug/L	97	2.4	SW846 8260B
Chloromethane	10	7.8	ug/L	78		SW846 8260B
	10	8.2	ug/L	82	4.1	SW846 8260B
1,2-Dibromo-3-chloropropane	10	7.6	ug/L	76		SW846 8260B
	10	7.8	ug/L	78	2.2	SW846 8260B
1,2-Dibromoethane	10	9.8	ug/L	98		SW846 8260B
	10	10	ug/L	100	1.9	SW846 8260B
1,1-Dichloroethane	10	9.8	ug/L	98		SW846 8260B
	10	10	ug/L	100	1.8	SW846 8260B
1,2-Dichloroethane	10	10	ug/L	101		SW846 8260B
	10	10	ug/L	101	0.18	SW846 8260B
cis-1,2-Dichloroethene	10	9.4	ug/L	94		SW846 8260B
	10	9.5	ug/L	95	1.2	SW846 8260B
trans-1,2-Dichloroethene	10	9.7	ug/L	97		SW846 8260B
	10	10	ug/L	102	5.3	SW846 8260B
1,1-Dichloroethene	10	10	ug/L	101		SW846 8260B
	10	10	ug/L	103	1.8	SW846 8260B
1,2-Dichloroethene (total)	20	19	ug/L	96		SW846 8260B
	20	20	ug/L	99	3.3	SW846 8260B
1,2-Dichloropropane	10	10	ug/L	100		SW846 8260B
	10	9.6	ug/L	96	3.4	SW846 8260B
cis-1,3-Dichloropropene	10	9.2	ug/L	92		SW846 8260B
	10	9.2	ug/L	92	0.13	SW846 8260B
trans-1,3-Dichloropropene	10	8.5	ug/L	85		SW846 8260B
	10	8.4	ug/L	84	1.0	SW846 8260B

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LABORATORY CONTROL SAMPLE DATA REPORT

GC/MS Volatiles

PARAMETER	SPIKE	MEASURED		PERCENT		METHOD
	AMOUNT	AMOUNT	UNITS	RECOVERY	RPD	
Ethylbenzene	10	10	ug/L	100		SW846 8260B
	10	10	ug/L	100	0.71	SW846 8260B
2-Hexanone	10	8.1	ug/L	81		SW846 8260B
	10	8.6	ug/L	86	6.0	SW846 8260B
Methylene chloride	10	13 a	ug/L	133		SW846 8260B
	10	14 a	ug/L	138	3.8	SW846 8260B
4-Methyl-2-pentanone	10	11	ug/L	107		SW846 8260B
	10	12	ug/L	117	8.9	SW846 8260B
Styrene	10	11	ug/L	110		SW846 8260B
	10	11	ug/L	108	1.8	SW846 8260B
1,1,2,2-Tetrachloroethane	10	11	ug/L	113		SW846 8260B
	10	12	ug/L	116	2.3	SW846 8260B
Tetrachloroethene	10	9.6	ug/L	96		SW846 8260B
	10	9.8	ug/L	98	1.6	SW846 8260B
Toluene	10	9.5	ug/L	95		SW846 8260B
	10	9.4	ug/L	94	1.2	SW846 8260B
1,1,1-Trichloroethane	10	8.8	ug/L	88		SW846 8260B
	10	8.7	ug/L	87	1.4	SW846 8260B
1,1,2-Trichloroethane	10	9.8	ug/L	98		SW846 8260B
	10	9.9	ug/L	99	1.3	SW846 8260B
Trichloroethene	10	9.6	ug/L	96		SW846 8260B
	10	9.6	ug/L	96	0.010	SW846 8260B
Trichlorofluoromethane	10	8.0	ug/L	80		SW846 8260B
	10	8.2	ug/L	82	1.8	SW846 8260B
Vinyl chloride	10	8.0	ug/L	80		SW846 8260B
	10	8.2	ug/L	82	2.5	SW846 8260B
Xylenes (total)	30	32	ug/L	106		SW846 8260B
	30	31	ug/L	105	0.96	SW846 8260B

<u>SURROGATE</u>	PERCENT <u>RECOVERY</u>	RECOVERY <u>LIMITS</u>
Dibromofluoromethane	104	(73 - 122)
1,2-Dichloroethane-d4	104	(73 - 122)
Toluene-d8	104	(61 - 128)
4-Bromofluorobenzene	98	(76 - 110)
	97	(76 - 110)
	108	(74 - 116)
	109	(74 - 116)

NOTE (S) :

Calculations are performed before rounding to avoid round-off errors in calculated results.

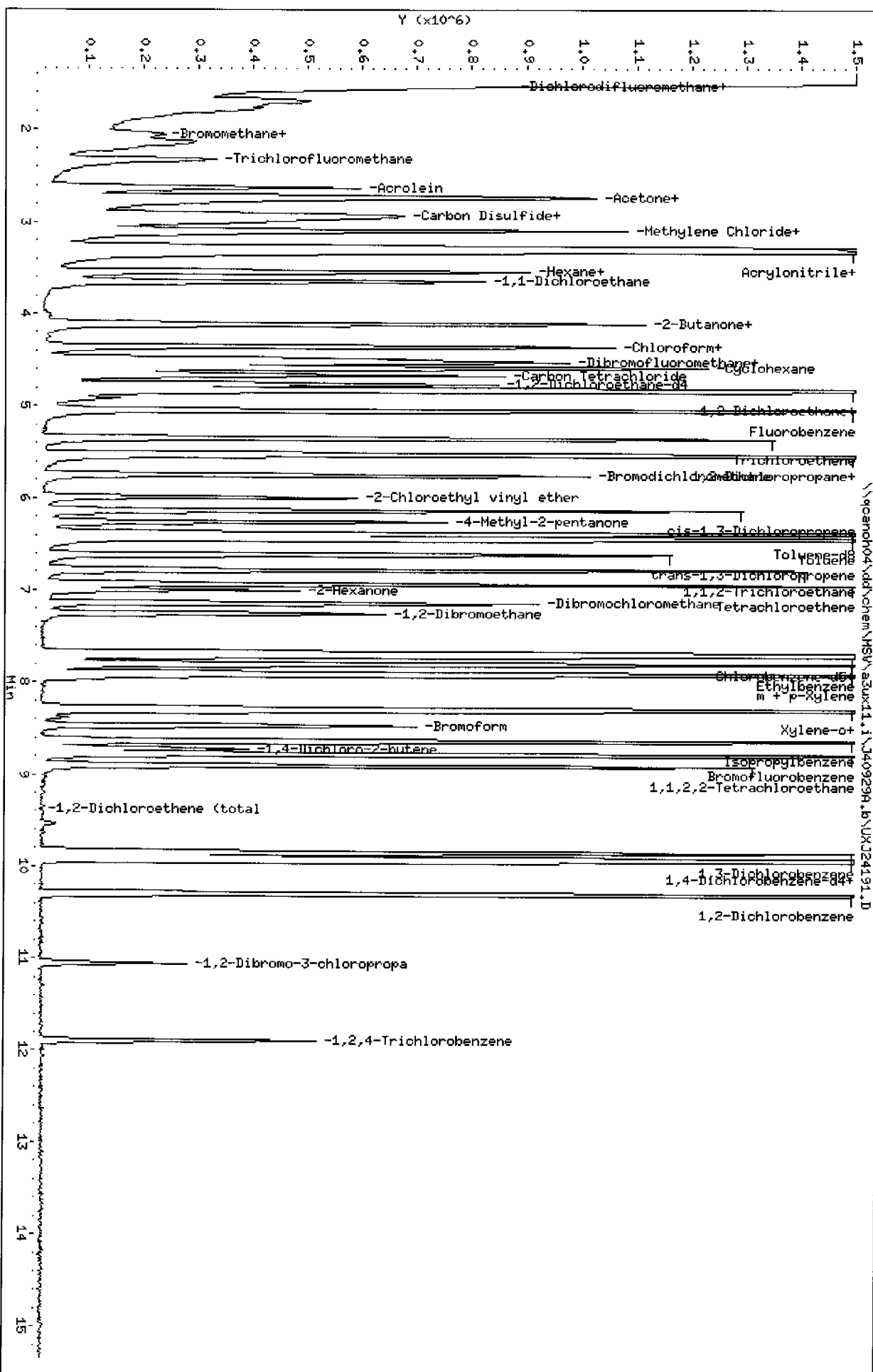
Bold print denotes control parameters

a Spiked analyte recovery is outside stated control limits.

Client ID: CHECK
 Sample Info: CHECK
 Purge Volume: 5.0
 Column phase: DB624

GRDLRIFAC

Instrument: 3ux11.i
 Operator: 43532
 Column diameter: 0.18



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24191.D
Report Date: 30-Sep-2004 08:37

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24191.D
Lab Smp Id: GRDLR1AC
Inj Date : 29-SEP-2004 08:37
Operator : 43582 Inst ID: a3ux11.i
Smp Info : CHECK
Misc Info : J40929A,8260LLUX11,2-8260.SUB,43582,3
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 3 QC Sample: METHSPIKE
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ug/L)
*	1 Fluorobenzene	96	5.041	5.041 (1.000)	2102753	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1771165	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	886729	50.0000		
\$	4 Dibromofluoromethane	113	4.485	4.473 (0.890)	510952	52.1903	10.438	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	698297	51.8707	10.374	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	2088903	49.0924	9.818	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	980182	54.2157	10.843	
8	Dichlorodifluoromethane	85	1.538	1.538 (0.305)	352669	32.0808	6.416	
9	Chloromethane	50	1.680	1.692 (0.333)	768480	39.1525	7.830	
10	Vinyl Chloride	62	1.775	1.775 (0.352)	539791	40.2043	8.041	
11	Bromomethane	94	2.059	2.047 (0.409)	231527	36.3563	7.271	
12	Chloroethane	64	2.142	2.130 (0.425)	452167	45.8249	9.165	
13	Trichlorofluoromethane	101	2.319	2.319 (0.460)	565299	40.1705	8.034	
15	Acrolein	56	2.627	2.627 (0.521)	841405	640.589	128.12	
16	Acetone	43	2.745	2.734 (0.545)	224482	44.6743	8.935	
17	1,1-Dichloroethene	96	2.722	2.722 (0.540)	475845	50.6022	10.120	
18	Freon-113	151	2.745	2.745 (0.545)	367336	62.0293	12.406	
19	Iodomethane	142		Compound Not Detected.				

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng)
20 Carbon Disulfide	76	2.911	2.911 (0.578)	1729637	49.2404	9.848	
21 Methylene Chloride	84	3.089	3.089 (0.613)	847733	66.4314	13.286	
22 Acetonitrile	41	2.958	2.947 (0.587)	753290	603.606	120.72	
23 Acrylonitrile	53	3.266	3.266 (0.648)	2143962	552.449	110.49	
24 Methyl tert-butyl ether	73	3.313	3.313 (0.657)	1291737	42.8471	8.569	
25 trans-1,2-Dichloroethene	96	3.325	3.313 (0.660)	541859	48.5735	9.715	
26 Hexane	86	3.538	3.538 (0.702)	98076	54.1706	10.834	
27 Vinyl acetate	43	3.538	3.668 (0.702)	319314	17.8881	3.578	
28 1,1-Dichloroethane	63	3.645	3.645 (0.723)	995488	48.9184	9.784	
29 tert-Butyl Alcohol	59	3.313	3.160 (0.657)	30049	36.7402	7.348	
30 2-Butanone	43	4.094	4.094 (0.812)	220126	38.2201	7.644	
M 31 1,2-Dichloroethene (total)	96			1094241	95.7015	19.140	
32 cis-1,2-dichloroethene	96	4.106	4.106 (0.815)	552382	47.1280	9.426	
33 2,2-Dichloropropane	77		Compound Not Detected.				
34 Bromochloromethane	128		Compound Not Detected.				
35 Chloroform	83	4.355	4.355 (0.864)	1016223	49.6762	9.935	
36 Tetrahydrofuran	42	4.343	4.343 (0.862)	11404	3.71219	0.7424	
37 1,1,1-Trichloroethane	97	4.520	4.520 (0.897)	677674	43.8629	8.772	
38 1,1-Dichloropropene	75		Compound Not Detected.				
39 Carbon Tetrachloride	117	4.662	4.662 (0.925)	571738	47.1306	9.426	
40 1,2-Dichloroethane	62	4.816	4.816 (0.955)	844839	50.5450	10.109	
41 Benzene	78	4.816	4.816 (0.955)	2403971	49.6565	9.931	
42 Trichloroethene	130	5.349	5.349 (1.061)	528743	47.8550	9.571	
43 1,2-Dichloropropene	63	5.526	5.526 (1.096)	596082	49.9133	9.983	
44 1,4-Dioxane	88		Compound Not Detected.				
45 Dibromomethane	93		Compound Not Detected.				
46 Bromodichloromethane	83	5.751	5.751 (1.141)	791552	50.3857	10.077	
47 2-Chloroethyl vinyl ether	63	5.988	5.988 (1.188)	287668	42.8401	8.568	
48 cis-1,3-Dichloropropene	75	6.130	6.130 (1.216)	881828	45.8768	9.175	
49 4-Methyl-2-pentanone	43	6.260	6.248 (1.242)	552771	53.4817	10.696	
50 Toluene	91	6.437	6.437 (0.838)	2510873	47.4676	9.494	
51 trans-1,3-Dichloropropene	75	6.603	6.603 (0.860)	805847	42.4906	8.498	
52 Ethyl Methacrylate	69		Compound Not Detected.				
53 1,1,2-Trichloroethane	97	6.780	6.769 (0.883)	523226	48.7649	9.753	
54 1,3-Dichloropropane	76		Compound Not Detected.				
55 Tetrachloroethene	164	6.934	6.934 (0.903)	411712	48.0416	9.608	
56 2-Hexanone	43	6.993	6.993 (0.911)	348699	40.7116	8.142	
57 Dibromochloromethane	129	7.135	7.135 (0.929)	575105	50.2530	10.051	
58 1,2-Dibromoethane	107	7.254	7.254 (0.945)	519459	49.1372	9.827	
59 Chlorobenzene	112	7.703	7.703 (1.003)	1748877	51.0622	10.212	
60 1,1,1,2-Tetrachloroethane	131		Compound Not Detected.				
61 Ethylbenzene	106	7.798	7.798 (1.015)	868170	50.1942	10.039	
62 m + p-Xylene	106	7.905	7.905 (1.029)	2373223	106.532	21.306	
M 63 Xylenes (total)	106			3516826	158.824	31.765	
64 Xylene-o	106	8.283	8.283 (1.079)	1143603	52.2917	10.458	
65 Styrene	104	8.295	8.295 (1.080)	2144282	54.8287	10.966	
66 Bromoform	173	8.473	8.473 (1.103)	412125	52.9800	10.596	

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
67 Isopropylbenzene	====	105	8.626	8.626 (1.123)	2666916	54.7531	10.951
68 1,1,2,2-Tetrachloroethane		83	8.899	8.899 (0.898)	756749	56.6143	11.323
69 1,4-Dichloro-2-butene		53	8.721	8.958 (0.881)	9719	2.20988	0.4420
70 1,2,3-Trichloropropane		110		Compound Not Detected.			
71 Bromobenzene		156		Compound Not Detected.			
72 n-Propylbenzene		120		Compound Not Detected.			
73 2-Chlorotoluene		126		Compound Not Detected.			
74 1,3,5-Trimethylbenzene		105		Compound Not Detected.			
75 4-Chlorotoluene		126		Compound Not Detected.			
76 tert-Butylbenzene		119		Compound Not Detected.			
77 1,2,4-Trimethylbenzene		105		Compound Not Detected.			
78 sec-Butylbenzene		105		Compound Not Detected.			
79 4-Isopropyltoluene		119		Compound Not Detected.			
80 1,3-Dichlorobenzene		146	9.845	9.845 (0.994)	1196027	48.9683	9.794
81 1,4-Dichlorobenzene		146	9.928	9.928 (1.002)	1319140	51.6226	10.324
82 n-Butylbenzene		91		Compound Not Detected.			
83 1,2-Dichlorobenzene		146	10.295	10.295 (1.039)	1175167	48.8502	9.770
84 1,2-Dibromo-3-chloropropane		157	11.064	11.052 (1.117)	86494	38.2123	7.642
85 1,2,4-Trichlorobenzene		180	11.892	11.892 (1.201)	183131	17.8809	3.576
86 Hexachlorobutadiene		225		Compound Not Detected.			
87 Naphthalene		128		Compound Not Detected.			
88 1,2,3-Trichlorobenzene		180		Compound Not Detected.			
98 Cyclohexane		56	4.580	4.580 (0.908)	682995	44.1510	8.830
143 Methyl Acetate		43	3.006	3.006 (0.596)	411507	47.5741	9.515
144 Methylcyclohexane		83	5.526	5.514 (1.096)	564425	43.4795	8.696
141 1,3,5-Trichlorobenzene		180		Compound Not Detected.			

Data File: \\\wparoh04\dd\chem\MSV\3Jux11.i\J40929A.b\UXJ24192.D
Date : 29-SEP-2004 09:00

Client ID:

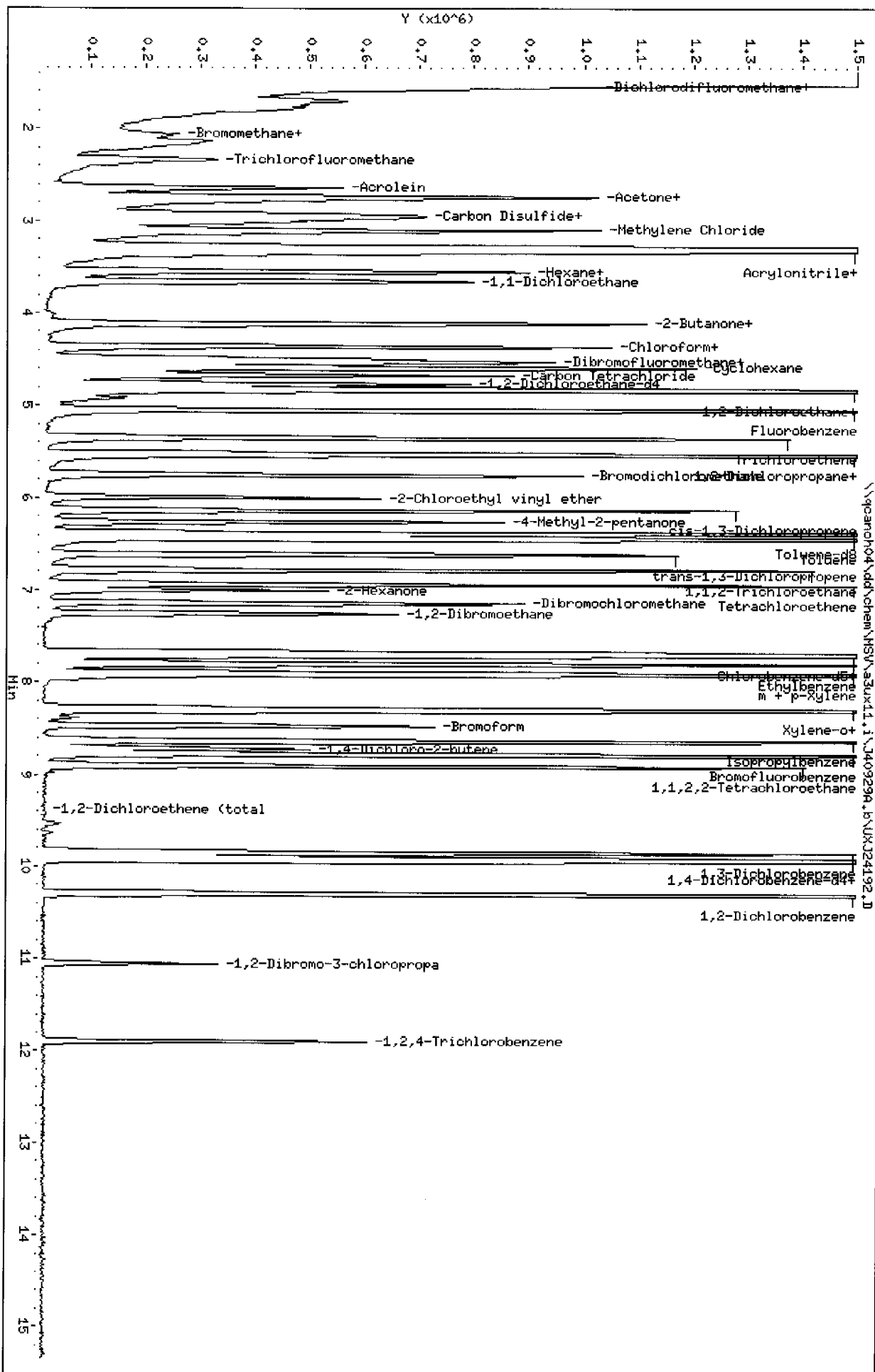
Client ID:

Sample Info CHECK

Column phase: DB624

o
624

Instrument: a3ux1.i
Operator: 43582
Column diameter: 0.18



STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24192.D
Lab Smp Id: grdlrlad
Inj Date : 29-SEP-2004 09:00
Operator : 43582 Inst ID: a3ux11.i
Smp Info : CHECK
Misc Info : J40929A, 8260LLUX11, 2-8260.SUB, 43582, 3
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 4 QC Sample: METHSPIKE
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 2-8260.SUB
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ug/L)
*	1 Fluorobenzene	96	5.041	5.041 (1.000)	2090526	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1767481	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	932881	50.0000		
\$	4 Dibromofluoromethane	113	4.485	4.473 (0.890)	507988	52.1910	10.438	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	696071	52.0078	10.402	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	2058681	48.4830	9.697	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	981509	54.4022	10.880	
8	Dichlorodifluoromethane	85	1.550	1.538 (0.308)	376522	34.4509	6.890	
9	Chloromethane	50	1.692	1.692 (0.336)	795945	40.7889	8.158	
10	Vinyl Chloride	62	1.799	1.775 (0.357)	550496	41.2414	8.248	
11	Bromomethane	94	2.059	2.047 (0.409)	245884	38.8366	7.767	
12	Chloroethane	64	2.130	2.130 (0.423)	470690	47.9811	9.596	
13	Trichlorofluoromethane	101	2.331	2.319 (0.463)	572135	40.8940	8.179	
15	Acrolein	56	2.639	2.627 (0.524)	864058	661.683	132.34	
16	Acetone	43	2.745	2.734 (0.545)	227320	45.6102	9.122	
17	1,1-Dichloroethene	96	2.734	2.722 (0.542)	481773	51.5322	10.306	
18	Freon-113	151	2.757	2.745 (0.547)	371526	63.0693	12.614	
19	Iodomethane	142		Compound Not Detected.				

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
20 Carbon Disulfide	76	2.923	2.911 (0.580)	1816264	52.0090	10.402	
21 Methylene Chloride	84	3.100	3.089 (0.615)	871461	69.0075	13.801	
22 Acetonitrile	41	2.958	2.947 (0.587)	892666	719.470	143.89	
23 Acrylonitrile	53	3.278	3.266 (0.650)	2218233	574.930	114.99	
24 Methyl tert-butyl ether	73	3.325	3.313 (0.660)	1239841	41.3663	8.273	
25 trans-1,2-Dichloroethene	96	3.325	3.313 (0.660)	568151	51.2283	10.246	
26 Hexane	86	3.550	3.538 (0.704)	104962	58.1703	11.634	
27 Vinyl acetate	43	3.550	3.668 (0.704)	329569	18.5705	3.714	
28 1,1-Dichloroethane	63	3.645	3.645 (0.723)	1008284	49.8370	9.967	
29 <i>tert</i> -Butyl Alcohol	59	3.006	3.160 (0.596)	34280	42.1585	8.432	
30 2-Butanone	43	4.106	4.094 (0.815)	247721	43.2630	8.652	
M 31 1,2-Dichloroethene (total)	96				1124169	98.9439	19.789
32 cis-1,2-dichloroethene	96	4.106	4.106 (0.815)	556018	47.7156	9.543	
33 2,2-Dichloropropane	77		Compound Not Detected.				
34 Bromochloromethane	128		Compound Not Detected.				
35 Chloroform	83	4.355	4.355 (0.864)	986837	48.5219	9.704	
36 Tetrahydrofuran	42	4.343	4.343 (0.862)	8586	2.86424	0.5728	
37 1,1,1-Trichloroethane	97	4.520	4.520 (0.897)	664548	43.2649	8.653	
38 1,1-Dichloropropene	75		Compound Not Detected.				
39 Carbon Tetrachloride	117	4.662	4.662 (0.925)	576326	47.7866	9.557	
40 1,2-Dichloroethane	62	4.828	4.816 (0.958)	838375	50.4517	10.090	
41 Benzene	78	4.828	4.816 (0.958)	2345068	48.7232	9.745	
42 Trichloroethene	130	5.349	5.349 (1.061)	525593	47.8482	9.570	
43 1,2-Dichloropropane	63	5.526	5.526 (1.096)	572786	48.2432	9.649	
44 1,4-Dioxane	88		Compound Not Detected.				
45 Dibromomethane	93		Compound Not Detected.				
46 Bromodichloromethane	83	5.751	5.751 (1.141)	775945	49.6811	9.936	
47 2-Chloroethyl vinyl ether	63	5.999	5.988 (1.190)	309719	46.3937	9.279	
48 cis-1,3-Dichloropropene	75	6.130	6.130 (1.216)	877798	45.9342	9.187	
49 4-Methyl-2-pentanone	43	6.260	6.248 (1.242)	601007	58.4887	11.698	
50 Toluene	91	6.437	6.437 (0.838)	2474694	46.8811	9.376	
51 trans-1,3-Dichloropropene	75	6.615	6.603 (0.861)	795769	42.0467	8.409	
52 Ethyl Methacrylate	69		Compound Not Detected.				
53 1,1,2-Trichloroethane	97	6.780	6.769 (0.883)	528976	49.4036	9.881	
54 1,3-Dichloropropane	76		Compound Not Detected.				
55 Tetrachloroethene	164	6.934	6.934 (0.903)	417496	48.8180	9.764	
56 2-Hexanone	43	6.993	6.993 (0.911)	369311	43.2080	8.642	
57 Dibromochloromethane	129	7.135	7.135 (0.929)	566653	49.6177	9.924	
58 1,2-Dibromoethane	107	7.254	7.254 (0.945)	528192	50.0674	10.013	
59 Chlorobenzene	112	7.703	7.703 (1.003)	1700618	49.7566	9.951	
60 1,1,1,2-Tetrachloroethane	131		Compound Not Detected.				
61 Ethylbenzene	106	7.798	7.798 (1.015)	860288	49.8421	9.968	
62 m + p-Xylene	106	7.905	7.905 (1.029)	2341167	105.312	21.062	
M 63 Xylenes (total)	106			3475818	157.303	31.460	
64 Xylene-o	106	8.283	8.283 (1.079)	1134651	51.9905	10.398	
65 Styrene	104	8.295	8.295 (1.080)	2100674	53.8256	10.765	
66 Bromoform	173	8.473	8.473 (1.103)	425809	54.8532	10.971	

Compounds	QUANT SIG	CONCENTRATIONS					
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)
67 Isopropylbenzene	105		8.638	8.626 (1.125)		2692750	55.3987
68 1,1,2,2-Tetrachloroethane	83		8.898	8.899 (0.898)		814941	57.9516
69 1,4-Dichloro-2-butene	53		8.721	8.958 (0.881)		13980	3.02147
70 1,2,3-Trichloropropane	110			Compound Not Detected.			
71 Bromobenzene	156			Compound Not Detected.			
72 n-Propylbenzene	120			Compound Not Detected.			
73 2-Chlorotoluene	126			Compound Not Detected.			
74 1,3,5-Trimethylbenzene	105			Compound Not Detected.			
75 4-Chlorotoluene	126			Compound Not Detected.			
76 tert-Butylbenzene	119			Compound Not Detected.			
77 1,2,4-Trimethylbenzene	105			Compound Not Detected.			
78 sec-Butylbenzene	105			Compound Not Detected.			
79 4-Isopropyltoluene	119			Compound Not Detected.			
80 1,3-Dichlorobenzene	146	9.845	9.845 (0.994)		1203365	46.8313	9.366
81 1,4-Dichlorobenzene	146	9.928	9.928 (1.002)		1349795	50.2090	10.042
82 n-Butylbenzene	91			Compound Not Detected.			
83 1,2-Dichlorobenzene	146	10.295	10.295 (1.039)		1192047	47.1004	9.420
84 1,2-Dibromo-3-chloropropane	157	11.052	11.052 (1.116)		93024	39.0640	7.813
85 1,2,4-Trichlorobenzene	180	11.892	11.892 (1.201)		214845	19.9397	3.988
86 Hexachlorobutadiene	225			Compound Not Detected.			
87 Naphthalene	128			Compound Not Detected.			
88 1,2,3-Trichlorobenzene	180			Compound Not Detected.			
98 Cyclohexane	56	4.591	4.580 (0.911)		706349	45.9278	9.186
143 Methyl Acetate	43	3.018	3.006 (0.599)		430160	50.0214	10.004
144 Methylcyclohexane	83	5.526	5.514 (1.096)		578857	44.8521	8.970
141 1,3,5-Trichlorobenzene	180			Compound Not Detected.			

METHOD BLANK REPORT

GC/MS Volatiles

Client Lot #....: A4I270179 **Work Order #....:** GRDLR1AA **Matrix.....:** WATER
MB Lot-Sample #: A4I290000-268
Analysis Date...: 09/29/04 **Prep Date.....:** 09/29/04 **Final Wgt/Vol..:** 5 mL
Dilution Factor: 1 **Prep Batch #....:** 4273268
Initial Wgt/Vol: 5 mL

PARAMETER	RESULT	REPORTING		
		LIMIT	UNITS	METHOD
Acetone	ND	10	ug/L	SW846 8260B
Acetonitrile	ND	20	ug/L	SW846 8260B
Acrolein	ND	20	ug/L	SW846 8260B
Acrylonitrile	ND	20	ug/L	SW846 8260B
Benzene	ND	1.0	ug/L	SW846 8260B
Bromodichloromethane	ND	1.0	ug/L	SW846 8260B
Bromoform	ND	1.0	ug/L	SW846 8260B
Bromomethane	ND	1.0	ug/L	SW846 8260B
2-Butanone	ND	10	ug/L	SW846 8260B
Carbon disulfide	ND	1.0	ug/L	SW846 8260B
Carbon tetrachloride	ND	1.0	ug/L	SW846 8260B
Chlorobenzene	ND	1.0	ug/L	SW846 8260B
Chloroprene	ND	2.0	ug/L	SW846 8260B
Dibromochloromethane	ND	1.0	ug/L	SW846 8260B
Chloroethane	ND	1.0	ug/L	SW846 8260B
Chloroform	ND	1.0	ug/L	SW846 8260B
Chloromethane	0.53 J	1.0	ug/L	SW846 8260B
3-Chloropropene	ND	2.0	ug/L	SW846 8260B
1,2-Dibromo-3-chloropropane (DBCP)	ND	2.0	ug/L	SW846 8260B
1,2-Dibromoethane	ND	1.0	ug/L	SW846 8260B
Dibromomethane	ND	1.0	ug/L	SW846 8260B
trans-1,4-Dichloro-2-butene	ND	1.0	ug/L	SW846 8260B
1,1-Dichloroethane	ND	1.0	ug/L	SW846 8260B
1,2-Dichloroethane	ND	1.0	ug/L	SW846 8260B
cis-1,2-Dichloroethene	ND	1.0	ug/L	SW846 8260B
trans-1,2-Dichloroethene	ND	1.0	ug/L	SW846 8260B
1,1-Dichloroethene	ND	1.0	ug/L	SW846 8260B
1,2-Dichloroethene (total)	ND	2.0	ug/L	SW846 8260B
Dichlorofluoromethane	ND	2.0	ug/L	SW846 8260B
1,2-Dichloropropane	ND	1.0	ug/L	SW846 8260B
cis-1,3-Dichloropropene	ND	1.0	ug/L	SW846 8260B
trans-1,3-Dichloropropene	ND	1.0	ug/L	SW846 8260B
1,4-Dioxane	ND	50	ug/L	SW846 8260B
Ethylbenzene	ND	1.0	ug/L	SW846 8260B
Ethyl methacrylate	ND	1.0	ug/L	SW846 8260B
2-Hexanone	ND	10	ug/L	SW846 8260B
Iodomethane	ND	1.0	ug/L	SW846 8260B
Isobutanol	ND	50	ug/L	SW846 8260B

(Continued on next page)

METHOD BLANK REPORT

GC/MS Volatiles

Client Lot #....: A4I270179

Work Order #....: GRDLR1AA

Matrix.....: WATER

<u>PARAMETER</u>	<u>RESULT</u>	<u>REPORTING LIMIT</u>	<u>UNITS</u>	<u>METHOD</u>
Methacrylonitrile	ND	2.0	ug/L	SW846 8260B
Methylene chloride	ND	1.0	ug/L	SW846 8260B
Methyl methacrylate	ND	2.0	ug/L	SW846 8260B
4-Methyl-2-pentanone	ND	10	ug/L	SW846 8260B
Propionitrile	ND	4.0	ug/L	SW846 8260B
Styrene	ND	1.0	ug/L	SW846 8260B
1,1,1,2-Tetrachloroethane	ND	1.0	ug/L	SW846 8260B
1,1,2,2-Tetrachloroethane	ND	1.0	ug/L	SW846 8260B
Tetrachloroethene	ND	1.0	ug/L	SW846 8260B
Toluene	ND	1.0	ug/L	SW846 8260B
1,1,1-Trichloroethane	ND	1.0	ug/L	SW846 8260B
1,1,2-Trichloroethane	ND	1.0	ug/L	SW846 8260B
Trichloroethene	ND	1.0	ug/L	SW846 8260B
Trichlorofluoromethane	ND	1.0	ug/L	SW846 8260B
1,2,3-Trichloropropane	ND	1.0	ug/L	SW846 8260B
Vinyl acetate	ND	2.0	ug/L	SW846 8260B
Vinyl chloride	ND	1.0	ug/L	SW846 8260B
Xylenes (total)	ND	2.0	ug/L	SW846 8260B

<u>SURROGATE</u>	<u>PERCENT RECOVERY</u>	<u>RECOVERY LIMITS</u>
Dibromofluoromethane	110	(73 - 122)
1,2-Dichloroethane-d4	107	(61 - 128)
Toluene-d8	89	(76 - 110)
4-Bromofluorobenzene	86	(74 - 116)

NOTE(S) :

Calculations are performed before rounding to avoid round-off errors in calculated results.

J Estimated result. Result is less than RL.

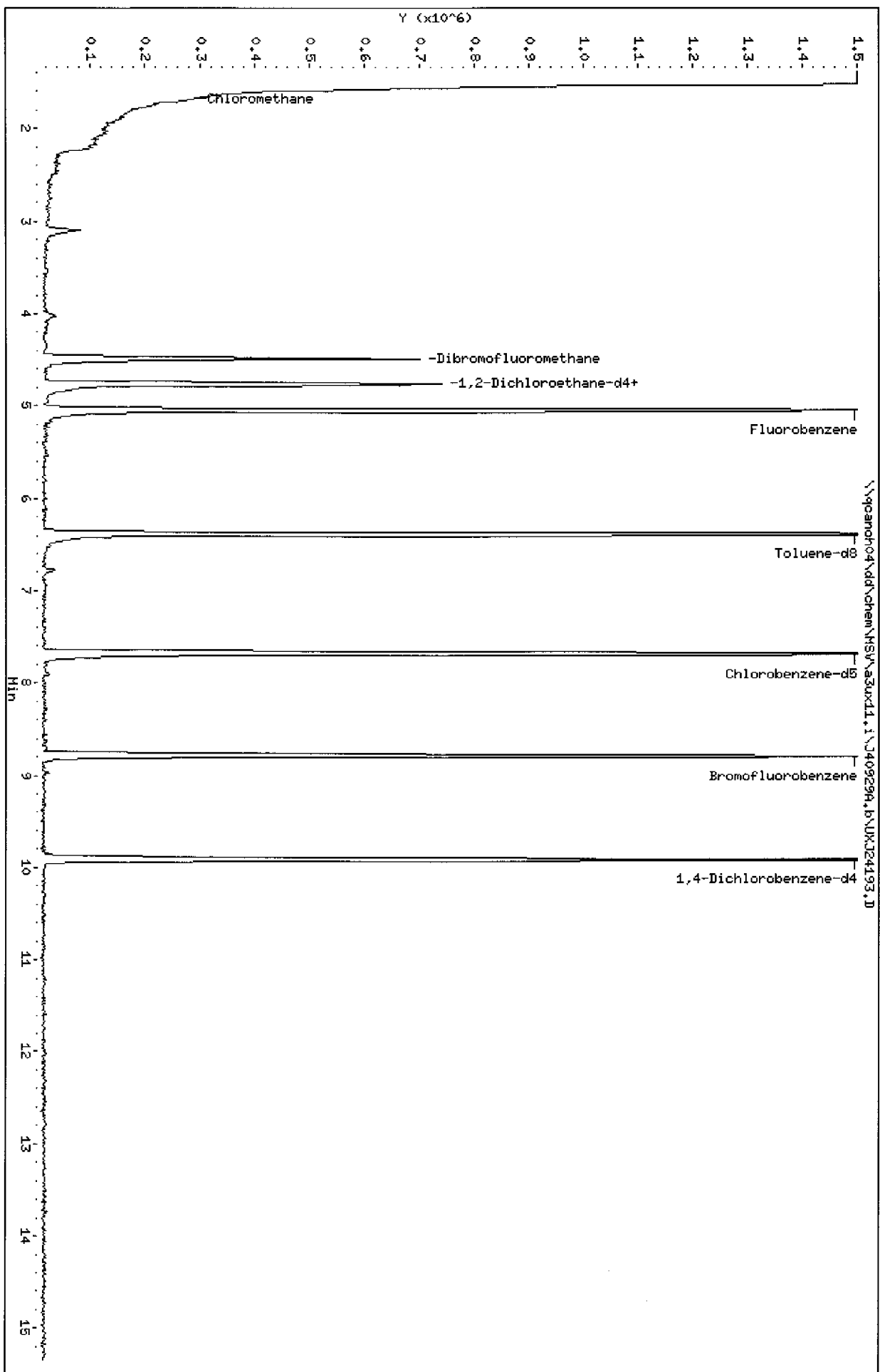
Data File: \\pcancho4\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UKJ24193.D
Date : 28-SEP-2004 09:22

Client ID:
Sample Info: VRLK
Purge Volume: 5.0
Column phase: DE624

GRDLR IAR

Instrument: a3ux11.i
Operator: 43582
Column diameter: 0.18

\\pcancho4\\dd\\chem\\MSV\\a3ux11.i\\J40929A.b\\UKJ24193.D



Data File: \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24193.D
Report Date: 30-Sep-2004 08:38

STL North Canton

VOLATILE REPORT SW-846 Method

Data file : \\qcanoh04\dd\chem\MSV\a3ux11.i\J40929A.b\UXJ24193.D
Lab Smp Id: grdlrlaa
Inj Date : 29-SEP-2004 09:22
Operator : 43582 Inst ID: a3ux11.i
Smp Info : VBLK
Misc Info : J40929A,8260LLUX11,,43582,3,,BLANK,,0
Comment :
Method : \\QCANOH04\dd\chem\MSV\a3ux11.i\J40929A.b\8260LLUX11.m
Meth Date : 30-Sep-2004 08:36 evansl Quant Type: ISTD
Cal Date : 14-SEP-2004 15:41 Cal File: UXJ23875.D
Als bottle: 5 QC Sample: BLANK
Dil Factor: 1.00000
Integrator: HP RTE Compound Sublist: 4-8260+IX.sub
Target Version: 4.04
Processing Host: CANPMSV07

Concentration Formula: Amt * DF * 1/Vo

Name	Value	Description
DF	1.000	Dilution Factor
Vo	5.000	Sample volume

Compounds	QUANT SIG	CONCENTRATIONS						
		MASS	RT	EXP RT	REL RT	RESPONSE	(ng)	(ug/L)
*	1 Fluorobenzene	96	5.041	5.041 (1.000)	1924922	50.0000		
*	2 Chlorobenzene-d5	117	7.680	7.680 (1.000)	1674688	50.0000		
*	3 1,4-Dichlorobenzene-d4	152	9.904	9.904 (1.000)	707660	50.0000		
\$	4 Dibromofluoromethane	113	4.485	4.473 (0.890)	493006	55.0094	11.002	
\$	5 1,2-Dichloroethane-d4	65	4.757	4.757 (0.944)	662309	53.7425	10.748	
\$	6 Toluene-d8	98	6.378	6.378 (0.831)	1792480	44.5529	8.910	
\$	7 Bromofluorobenzene	95	8.780	8.780 (1.143)	734712	42.9794	8.596	
8	Dichlorodifluoromethane	85	Compound Not Detected.					
9	Chloromethane	50	1.680	1.692 (0.333)	47333	2.63430	0.5269	
10	Vinyl Chloride	62	Compound Not Detected.					
11	Bromomethane	94	Compound Not Detected.					
12	Chloroethane	64	Compound Not Detected.					
13	Trichlorofluoromethane	101	Compound Not Detected.					
15	Acrolein	56	Compound Not Detected.					
16	Acetone	43	Compound Not Detected.					
17	1,1-Dichloroethene	96	Compound Not Detected.					
18	Freon-113	151	Compound Not Detected.					

Compounds	QUANT SIG	MASS	CONCENTRATIONS				
			RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ng) FINAL (ug/L)
19 Iodomethane		142				Compound Not Detected.	
20 Carbon Disulfide		76				Compound Not Detected.	
21 Methylene Chloride		84				Compound Not Detected.	
22 Acetonitrile		41				Compound Not Detected.	
23 Acrylonitrile		53				Compound Not Detected.	
24 Methyl tert-butyl ether		73				Compound Not Detected.	
25 trans-1,2-Dichloroethene		96				Compound Not Detected.	
26 Hexane		86				Compound Not Detected.	
27 Vinyl acetate		43				Compound Not Detected.	
28 1,1-Dichloroethane		63				Compound Not Detected.	
29 tert-Butyl Alcohol		59				Compound Not Detected.	
30 2-Butanone		43				Compound Not Detected.	
M 31 1,2-Dichloroethene (total)		96				Compound Not Detected.	
32 cis-1,2-dichloroethene		96				Compound Not Detected.	
33 2,2-Dichloropropane		77				Compound Not Detected.	
34 Bromochloromethane		128				Compound Not Detected.	
35 Chloroform		83				Compound Not Detected.	
36 Tetrahydrofuran		42				Compound Not Detected.	
37 1,1,1-Trichloroethane		97				Compound Not Detected.	
38 1,1-Dichloropropene		75				Compound Not Detected.	
39 Carbon Tetrachloride		117				Compound Not Detected.	
40 1,2-Dichloroethane		62				Compound Not Detected.	
41 Benzene		78	4.828	4.816 (0.958)		37742	0.85162 0.1703
42 Trichloroethene		130				Compound Not Detected.	
43 1,2-Dichloropropane		63				Compound Not Detected.	
44 1,4-Dioxane		88				Compound Not Detected.	
45 Dibromomethane		93				Compound Not Detected.	
46 Bromodichloromethane		83				Compound Not Detected.	
47 2-Chloroethyl vinyl ether		63				Compound Not Detected.	
48 cis-1,3-Dichloropropene		75				Compound Not Detected.	
49 4-Methyl-2-pentanone		43				Compound Not Detected.	
50 Toluene		91				Compound Not Detected.	
51 trans-1,3-Dichloropropene		75				Compound Not Detected.	
52 Ethyl Methacrylate		69				Compound Not Detected.	
53 1,1,2-Trichloroethane		97				Compound Not Detected.	
54 1,3-Dichloropropane		76				Compound Not Detected.	
55 Tetrachloroethene		164				Compound Not Detected.	
56 2-Hexanone		43				Compound Not Detected.	
57 Dibromochloromethane		129				Compound Not Detected.	
58 1,2-Dibromoethane		107				Compound Not Detected.	
59 Chlorobenzene		112				Compound Not Detected.	
60 1,1,1,2-Tetrachloroethane		131				Compound Not Detected.	
61 Ethylbenzene		106				Compound Not Detected.	
62 m + p-Xylene		106				Compound Not Detected.	
M 63 Xylenes (total)		106				Compound Not Detected.	
64 Xylene-o		106				Compound Not Detected.	
65 Styrene		104				Compound Not Detected.	

Compounds	QUANT SIG	MASS	CONCENTRATIONS					ON-COLUMN (ng)	FINAL (ug/L)
			RT	EXP RT	REL RT	RESPONSE	=====		
66 Bromoform		173				Compound Not Detected.			
67 Isopropylbenzene		105				Compound Not Detected.			
68 1,1,2,2-Tetrachloroethane		83				Compound Not Detected.			
69 1,4-Dichloro-2-butene		53				Compound Not Detected.			
70 1,2,3-Trichloropropane		110				Compound Not Detected.			
71 Bromobenzene		156				Compound Not Detected.			
72 n-Propylbenzene		120				Compound Not Detected.			
73 2-Chlorotoluene		126				Compound Not Detected.			
74 1,3,5-Trimethylbenzene		105				Compound Not Detected.			
75 4-Chlorotoluene		126				Compound Not Detected.			
76 tert-Butylbenzene		119				Compound Not Detected.			
77 1,2,4-Trimethylbenzene		105				Compound Not Detected.			
78 sec-Butylbenzene		105				Compound Not Detected.			
79 4-Isopropyltoluene		119				Compound Not Detected.			
80 1,3-Dichlorobenzene		146				Compound Not Detected.			
81 1,4-Dichlorobenzene		146				Compound Not Detected.			
82 n-Butylbenzene		91				Compound Not Detected.			
83 1,2-Dichlorobenzene		146				Compound Not Detected.			
84 1,2-Dibromo-3-chloropropane		157				Compound Not Detected.			
85 1,2,4-Trichlorobenzene		180				Compound Not Detected.			
86 Hexachlorobutadiene		225				Compound Not Detected.			
87 Naphthalene		128				Compound Not Detected.			
88 1,2,3-Trichlorobenzene		180				Compound Not Detected.			
14 Dichlorofluoromethane		67				Compound Not Detected.			
89 Ethyl Ether		59				Compound Not Detected.			
91 3-Chloropropene		76				Compound Not Detected.			
92 Isopropyl Ether		87				Compound Not Detected.			
93 2-Chloro-1,3-butadiene		53				Compound Not Detected.			
94 Propionitrile		54				Compound Not Detected.			
95 Ethyl Acetate		43				Compound Not Detected.			
96 Methacrylonitrile		41				Compound Not Detected.			
97 Isobutanol		41				Compound Not Detected.			
99 n-Butanol		56				Compound Not Detected.			
100 Methyl Methacrylate		41				Compound Not Detected.			
101 2-Nitropropane		41				Compound Not Detected.			
103 Cyclohexanone		55				Compound Not Detected.			
98 Cyclohexane		56				Compound Not Detected.			
143 Methyl Acetate		43				Compound Not Detected.			
144 Methylcyclohexane		83				Compound Not Detected.			
141 1,3,5-Trichlorobenzene		180				Compound Not Detected.			

Data File: \\qcanoh04\\dd\\chem\\MSV\\a3ux11.i\\340929A.b\\UXJ24193.D

Date : 29-SEP-2004 09:22

Client ID:

Instrument: a3ux11,i

Sample Info: VBLK

Purge Volume: 5.0

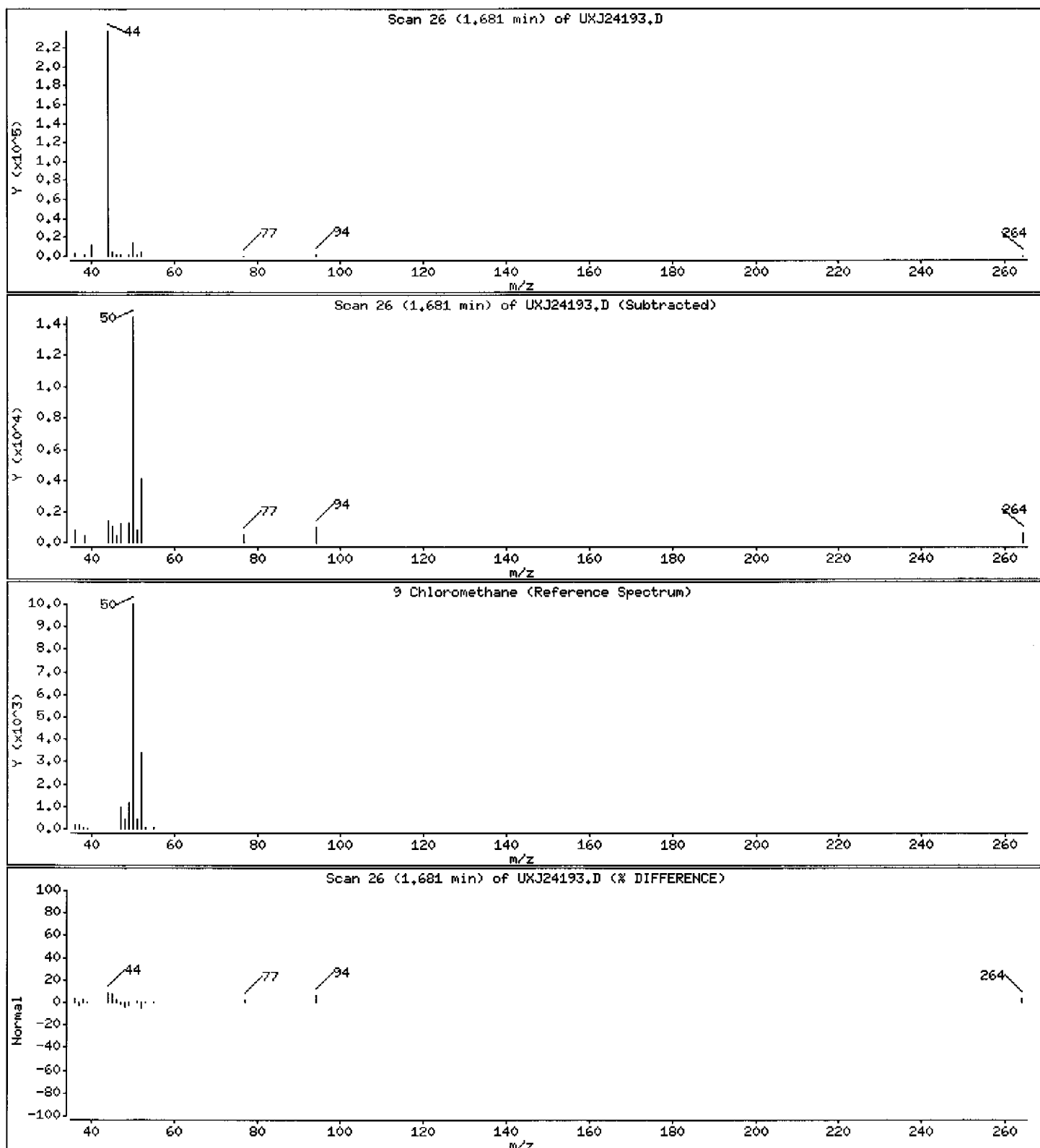
Operator: 43582

Column phase: DB624

Column diameter: 0.18

9 Chloromethane

Concentration: 0.5269 ug/L



Data File: \\qcanoh04\dd\chem\MSV\s3ux11.i\J40929A.b\UXJ24193.D

Date : 29-SEP-2004 09:22

Client ID:

Instrument: s3ux11.i

Sample Info: VBLK

Purge Volume: 5.0

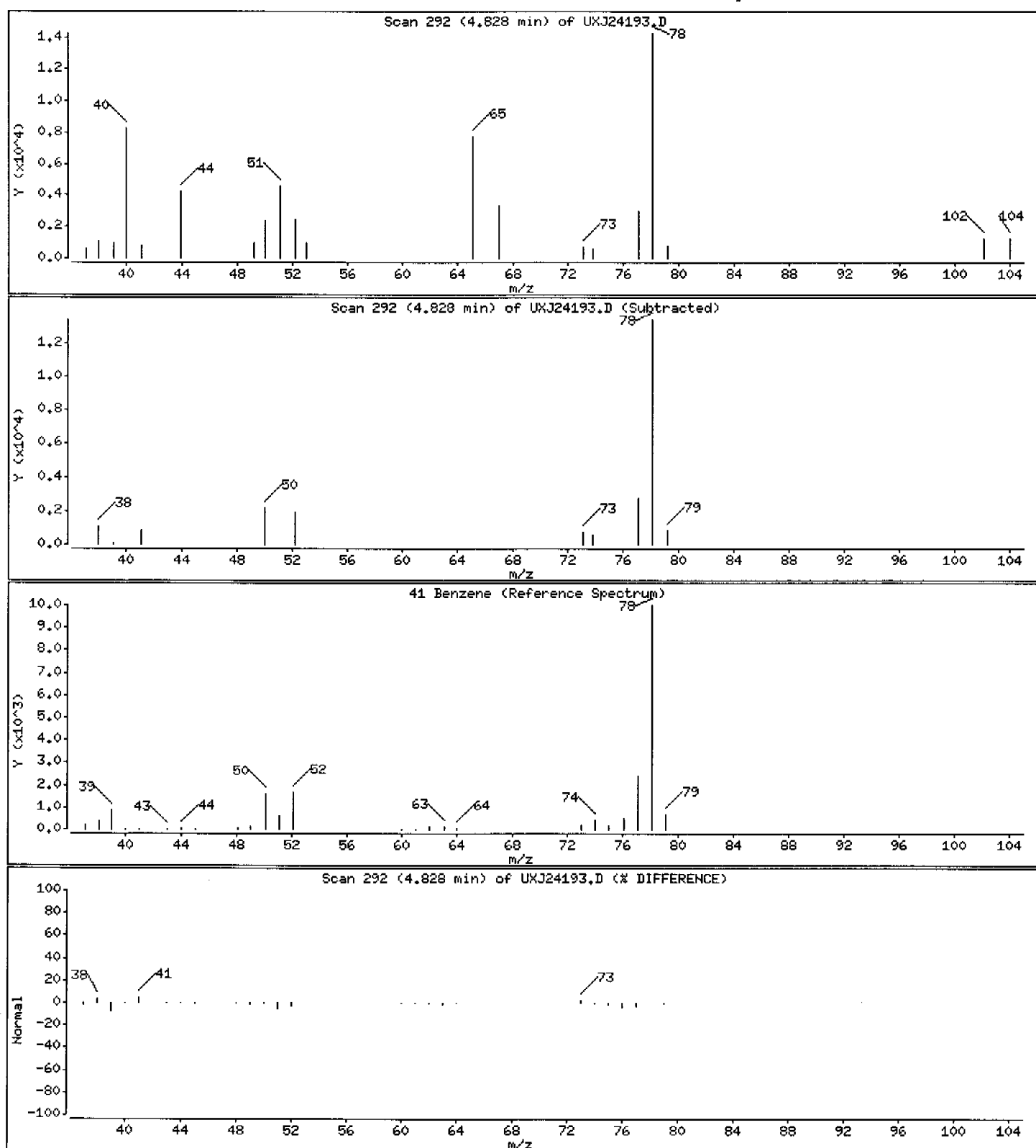
Operator: 43582

Column phase: DB624

Column diameter: 0.18

41 Benzene

Concentration: 0.1703 ug/L



**SEVERN
TRENT**

STL

MISCELLANEOUS DATA

UX11
Batch # _____

**STL-North Canton
GC/MS VOA Run Log**

Date: 8/16/04

Audrey

Analyst: _____
Level 2 review:

4

UX11
Batch # 4259349

STL-North Canton
GC/MS VOA Run Log

(9/16)

Date: 9/14/04

Column	BFB	Analysis	Purge & Trap
Type: DB624	100 C for 0.1 min	45 C for 2 min	Trap: #10
Length 20 M	to 200 C @ 20 C/min	to 200 C @ 20 C/min	Purge: 11
I.D. 0.18 mm	Hold 0 min	to 0 C @ 0 C/min	Desorb: 1 min @ 240 C
Flow Rate 0.4ml/min	IS # <u>V2246</u> SS # <u>V2247</u>	Hold 3 min	Bake: 5 min @ 250 C
			Heated purge: Yes No
v bpx	DBP 232	SONy	On
✓ 101 STD	UXR23870	200 K	On
✓ 101 STD	71	100 K	On
✓ 101 STD	72	SOMy	On
✓ 101 STD	73	25 K	On
✓ 101 STD	74	10 K	On
✓ 101 STD	75	5 K	On
✓ ACV	76	SONy	On
✓ Check GP86W	77	1	On
✓ Check Dup	78	1	On
✓ Blank	79	Sme	On
✓ GPMELIAA	80	0.2mle/ml	On
✓ GPMELIAA (S)	81	+SONy	On
✓ GPMELIAA (O)	82	—	On
✓ GPMELIAA ED	83	Sme	On
✓ GPHTDIAA	84	—	On
✓ GPJWVIAA ED	85	—	On
✓ GPJWVIAA	86	—	On
✓ GPJW3IAA	87	—	On
✓ GPJW4IAA	88	—	On
✓ GPJW5IAA	89	—	On
✓ GPJW6IAA	90	—	On
✓ GPJW7IAA	91	—	On
✓ GPJW8IAA	92	—	On
✓ GPJW9IAA	93	—	On
✓ GPKLEIAA	6m	84	On
✓ GPKLEIAA	—	85	On
✓ GPKMIAA	—	96	OK
✓ GPLDJIAA	m	97	On
✓ GPLDPAA	m	98	OK
✓ GPLOTIAA	—	99	On

Analyst: J. Boni
Level 2 review: J. Boni

29

UX11
Batch # 4273268

STL-North Canton
GC/MS VOA Run Log

9/89

Date: 9/29/04

Column	BFB	Analysis	Purge & Trap
Type: DB624	100 C for 0.1 min	45 C for 2 min	Trap: #10
Length 20 M	to 200 C @ 20 C/min	to 200 C @ 20 C/min	Purge: 11
I.D. 0.18 mm	Hold 0 min	to 0 C @ 0 C/min	Desorb: 1 min @ 240 C
Flow Rate 0.4ml/min	Hold 3 min	Hold 3 min	Bake: 5 min @ 250 C
IS # V2313	SS # V2314		Heated purge: Yes No
1 BFB /	1 BFB 243 song	dir inject (9:25) /	On
1/1 VOI STD	UKT0108	V230708/15 J40914	On
1/2 Ag STD	90	V2319 J40816	On
1/3 Check GROCR	91	V2302	On
1/4 Check Dup	92		On
1/5 Blank	93	Sm	On
1/6 GOKRMIAA	94	0.1me/Sm	Off
1/7 GOFERZIAZ	95	Sm	On
1/8 GOFER3IAZ	96	2AC(P) 3.5 me	On
1/9 GOFER4IAZ	97	2AC(P) 1.75 me	On
1/10 GOKRJIAA	98		On
1/11 GOKRLIAA	99		On
1/12 GOLDFIAA	100		On
1/13 GOLDFGIAA	01		On
1/14 GOKRMIAA	02	0.05me/Sm	On
1/15 GOG664IAA	03	1me/Sm	On
1/16 GOFER3ZAC	04	3.5me/Sm	On
1/17 GOFER4ZAC	05	1.75me/Sm	On
1/18 GONVPIAA	06	2me/Sm	Off me
1/19 GODJLIAA ED	07	2me/Sm	Off me
1/20 GQDQWIAA	08	0.005me/Sm	On
1/21 GONVLLIAA ED	09	Sm	On
1/22 GONVRIAA 1	10		On
1/23 GOKRMIAZ(S)	11	0.0me/Sm + song	On
1/24 GOKRMIAZ(O)	12		On
1/25 GONVLTIAA ED	13	Sm	On
1/26 GONVNTIAA 1	14		On
1/27 GQDQJQIAA ED	15		On
1/28 GQDQJHIAA	16		On
1/29 GOKR2IAA	17		On
1/30 GONVPIAA ED	18	1me/Sm	On
Analyst: JH			9/30/04
Level 2 review: SAR			JH

Severn Trent Laboratories, Inc

System Date: 9/27/04 10:56:33

Local Date: 9/27/04 12:56:33

MSVOC

Lot Summary - A4I270179

CLIENT: 5670 PAYNE FIRM INC.

SDG: 4I27179

Date Received: 9/25/04

PROJECT MANAGER: Roger K. Toth

Date Analysis Due: 9/29/04 N

SITE: EMD/OHIO

Date Report Due: 10/05/04

LOT COMMENTS:

Turnaround Time: 4

QC PACKAGE: Expanded Deliverables

FC = N

44

= Field(s) Changed

- - - - - R E P R I N T - - - - -

SAMP# W/O NO. PARAMETER X-REF Sampled Expires Est Sample ID, Comments / Analysis Comments

001- GQ67F-1AA XX I 25 QK 01 MS8260LL 9/24/04 10/08/04 Y #VE539/12.5-17.5/092404
 3 # # # # # Added 9#58 # Q: CLP MSVOA TCL Standard List
 EXP DEL, SDG# 4I27179 (CLSD), 8260 NEEDS TO HAVE 10X
 LESSER DIL, 48 HOUR TAT!
 AP9 Compounds

002- GQ67G-1AA XX I 25 QK 01 MS8260LL 9/24/04 10/08/04 Y #TRIP BLANK/092404
 2 # # # # # Added # # Q: CLP MSVOA TCL Standard List
 EXP DEL, SDG# 4I27179 (CLSD), 8260 NEEDS TO HAVE 10X
 LESSER DIL, 48 HOUR TAT!
 AP9 Compounds

PHz

AP

10/04/04 07:32:12 Sample Control Chain of Custody - STL North Canton PAGE 1

LOT NUMBER	SAMPLE ID	LAB	ANALYSIS TYPE	ANALYSIS DATE	ANALYST
A4127019	1	GG67F1AA	MS8260LL	9/29/04	Laura Evans
A4127019	2	GG67G1AA	MS8260LL	9/29/04	Laura Evans

* * * E N D O F R E P O R T * * *

END OF REPORT